## (19) World Intellectual Property Organization International Bureau





### (43) International Publication Date 30 August 2001 (30.08.2001)

**PCT** 

## (10) International Publication Number WO 01/63244 A1

- (51) International Patent Classification7: G01N 9/00, 33/48
- (21) International Application Number: PCT/US01/05150
- (22) International Filing Date: 16 February 2001 (16.02.2001)
- (25) Filing Language:

1.

English

(26) Publication Language:

English

(30) Priority Data:

09/514,026

25 February 2000 (25.02.2000) US

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- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



(54) Title: SOLUTION AND CRYSTAL STRUCTURES OF MMP-13 ACTIVE SITE AND USES THEREOF

(57) Abstract: The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), as well as to (i) methods of using the MMP-13 structure to rationally design or identify compounds or molecules that inhibit or activate MMP-13 activity, and (ii) compounds identified using said methods.

Docket No. 2368/16

# SOLUTION AND CRYSTAL STRUCTURES OF MMP-13 ACTIVE SITE AND USES THEREOF

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#### Field of the Invention

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), as well as to (i) methods of using the MMP-13 structure to rationally design or identify compounds or molecules that inhibit or activate MMP-13 activity, and (ii) compounds identified using said methods.

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#### Background of the Invention

Human collagenase-3 (MMP-13) is a member of the matrix metalloproteinase (MMP) family which includes the collagenases, stromelysins and gelatinases. The MMPs are involved in the degradation of the extracellular matrix and are associated with normal tissue remodeling processes such as pregnancy, wound healing, and angiogenesis. MMP expression and activity is highly controlled because of the degradative nature of these enzymes, where an apparent loss in MMP regulation results in the pathological destruction of connective tissue and the ensuing disease state. Accordingly, MMPs are a highly active set of targets for the design of therapeutic agents for the disease areas of arthritis and oncology (for reviews, see Woessner, J. F., FASEB 1991; Ries, C., and Petrides, E., Biol. Chem. Hoppe-Seyler 1995; Browner, M. F., Perspect. Drug Discovery Des. 1995; Morphy, et al., Curr. Med. Chem. 1995; and Zask, et al., Curr. Pharm. Des. 1996).

MMP-13 was identified on the basis of differential expression in normal breast tissues and in breast carcinoma. In addition, its expression has been reported in squamous cell carcinomas of the larynx, head and neck, in HCS-2/8 human chondrosarcoma cells, during fetal ossification, and in articular cartilage of arthritic patients.

There have been a number of X-ray and NMR structures solved for the catalytic domain of MMPs complexed with a variety of inhibitors (see e.g., Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Science 1994; Lovejoy, et al., Ann. N. Y. Acad. Sci. 1994; Lovejoy, et al.,

Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl. Acad. Sci. U.S.A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. USA 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr. 1996; Gooley, et al., J. Biomol. NMR 1996; Betz, et al., 5 Eur. J. Biochem. 1997; Gonnella, et al., Bioorg. Med. Chem. 1997; and Moy, et al., Biochemistry 1998). There is a close similarity in the overall threedimensional fold for these proteins consistent with the relatively high sequence homology (> 40%). Despite this similarity in the MMP structures, there is a distinct substrate specificity between these enzymes indicative of specific 10 biological roles for the various MMPs and a corresponding association with unique disease processes. One example of this potential specificity is the overexpression of MMP-13 in breast carcinoma and MMP-1 in papillary carcinomas. Therefore, the current paradigm in the development of MMP inhibitors is to design specificity into the structures of the small molecule instead of developing 15 a broad spectrum MMP inhibitor (Birkedal-Hansen, et al., Crit. Rev. Oral Biol. Med. 1993; and Rockwell, et al., J. Am. Chem. Soc. 1996). The rationale behind this approach is that an inhibitor specific for the MMP uniquely associated with a disease process may potentially minimize toxic side effects. Therefore, extensive structural information for the various MMPs is critical for a 20 structure-based approach in designing inhibitor selectivity (Birkedal-Hansen, et al., Crit. Rev. Oral Biol. Med. 1993; Rockwell, et al., J. Am. Chem. Soc. 1996; Ghose, et al., J. Am. Chem. Soc. 1995; Hajduk, et al., J. Am. Chem. Soc. 1997; and Olejniczak, et al., J. Am. Chem. Soc. 1997).

This concept has been facilitated by the extensive structural data available for the MMPs where a significant difference in the size and shape of the S1' pocket has been observed (Moy, et al., Biochemistry 1998; Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Ann. N.Y. Acad. Sci. 1994; Lovejoy, et al., Biochemistry 1994; Lovejoy, et al., Science 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl.

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Acad. Sci. U.S.A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. U.S.A. 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr. 1996; Gooley, et al., J. Biomol. NMR 1996; Betz, et al., Eur. J. Biochem. 1997; and Gonnella, et al., Bioorg. Med. Chem. 1997). This structural difference across the MMP family provides an obvious approach for designing specificity into potent MMP inhibitors by designing compounds that appropriately fill the available space in the S1' pocket while taking advantage of sequence differences. A number of examples have been previously reported using this approach where some selectivity between MMPs has been achieved by incorporating a biphenyl into the S1' pocket (see e.g., Hajduk, et al., J. Am. Chem. Soc. 1997; and Olejniczak, et al., J. Am. Chem. Soc. 1997).

The inventors have determined both the solution and crystal structures of MMP-13, and, using rational drug design methods, have designed a novel, potent inhibitor that is highly selective for MMP-13.

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#### Summary of the Invention

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13), and more specifically, to the crystal and solution structures of MMP-13 complexed with the inhibitor N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide (hereinafter referred to as "Compound A"), as determined using crystallography, spectroscopy and various computer modeling techniques. Particularly, the invention is directed to an MMP-13 active site comprised of the three dimensional structures of various binding pockets located both to the right (S1', S2', S3') and left (S1, S2, S3) of the catalytic zinc of MMP-13, and most particularly is directed to the three dimensional structure of an MMP-13 active site comprising the catalytic zinc and the S1' binding pocket, which is critical to the design and selection of inhibitors with increased potency and specificity for MMP-13, or conversely, for the design and selection of inhibitors of matrix metalloproteinases that are specific against MMP-13.

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Accordingly, the present invention discloses a solution comprising a biologically active catalytic fragment of human collagenase-3 (MMP-13) complexed with Compound A, as well as a crystallized catalytic fragment of MMP-13 complexed with Compound A. The three dimensional structure of the catalytic fragment of MMP-13 is provided by the relative atomic structural coordinates of Figure 4, as obtained from spectroscopy data, and Figure 5, as obtained from crystallography data. Also provided is an active site of MMP-13, characterized by a catalytic zinc, a beta strand, a Ca<sup>2+</sup> binding loop, an alpha helix and a random coil region, wherein the beta strand of said active site preferably comprises residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1, the Ca<sup>2+</sup> binding loop comprises residues F75, D76, G77, P78, and S79 according to Figure 1, the alpha helix comprises residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1, and the random coil region comprises residues P139, I140, and Y141 according to Figure 1. Said active site is further characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case, ± a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å.

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In an alternate embodiment of the invention, an active site of MMP-13 is characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å.

The solution or crystal structural coordinates of MMP-13 or portions thereof as provided by this invention may be stored in a

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machine-readable form on a machine-readable storage medium, e.g. a computer hard drive, diskette, DAT tape, etc., for display as a three-dimensional shape or for other uses involving computer-assisted manipulation of, or computation based on, the structural coordinates or the three-dimensional structures they define. By way of example, the data defining the three dimensional structure of MMP-13 or an MMP-13 complex of the present invention, or of a portion of MMP-13 or an MMP-13 complex as disclosed herein, may be stored in a machine-readable storage medium, and may be displayed as a graphical three-dimensional representation of the relevant structural coordinates, typically using a computer capable of reading the data from said storage medium and programmed with instructions for creating the representation from such data.

Accordingly, the present invention provides a machine, such as a computer, programmed in memory with the coordinates of the MMP-13 molecule or molecular complex, or portions thereof (such as, by way of example, the coordinates of the MMP-13 catalytic zinc with adjacent S1', S2' and/or S3' binding pockets), together with a program capable of converting the coordinates into a three dimensional graphical representation of the structural coordinates on a display connected to the machine. A machine having a memory containing such data aids in the rational design or selection of inhibitors or activators of MMP-13 activity, including the evaluation of ability of a particular chemical entity to favorably associate with MMP-13 or an MMP-13 complex as disclosed herein, as well as in the modeling of compounds, proteins, complexes, etc. related by structural or sequence homology to MMP-13.

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The present invention is additionally directed to a method of

determining the three dimensional structure of a molecule or molecular complex
whose structure is unknown, comprising the steps of first obtaining crystals or a
solution of the molecule or molecular complex whose structure is unknown, and
then generating X-ray diffraction data from the crystallized molecule or
molecular complex and/or generating NMR data from the solution of the

molecule or molecular complex. The generated diffraction or spectroscopy data
from the molecule or molecular complex can then be compared with the known

three dimensional structure of MMP-13 as disclosed herein, and the three dimensional structure of the unknown molecule or molecular complex conformed to the known MMP-13 structure using standard techniques such as molecular replacement analysis, 2D, 3D and 4D isotope filtering, editing and triple resonance NMR techniques, and computer homology modeling.

Alternatively, a three dimensional model of the unknown molecule may be generated by generating a sequence alignment between MMP-13 and the unknown molecule, based on any or all of amino acid sequence identity, secondary structure elements or tertiary folds, and then generating by computer modeling a three dimensional structure for the molecule using the three dimensional structure of, and sequence alignment with, MMP-13.

The present invention further provides a method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of using a three dimensional structure of MMP-13 as defined by the relative structural coordinates of amino acids encoding MMP-13 to design or select a potential inhibitor or activator, and synthesizing or obtaining said potential inhibitor or activator. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of an empty MMP-13 active site in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors. The method of the present invention is preferably used to design or select inhibitors of MMP-13 activity.

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Alternatively, the present invention provides a method for identifying a potential inhibitor or activator that is selective for one or more members of the matrix metalloproteinase family except MMP-13, comprising the steps of (i) using the three dimensional structures of MMP-13 and the desired target matrix metalloproteinase(s) as defined by the relative structural coordinates of amino acids encoding MMP-13 and the target matrix metalloproteinase(s) in order to design or select such a potential inhibitor or

activator, and (ii) synthesizing or obtaining said potential inhibitor or activator. In this case, the potential inhibitor or activator is designed to incorporate chemical or steric features favorable for association with an active site of the desired matrix metalloproteinase(s) and unfavorable for association with an MMP-13 active site, preferably where said active site comprises the MMP-13 S1' pocket. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of empty MMP-13/matrix metalloproteinase active sites in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors.

Also provided by the present invention are the inhibitors and activators designed or selected using the methods disclosed herein.

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#### **Brief Description of the Figures**

Figure 1 depicts the amino acid sequence encoding the catalytic fragment of human MMP-13.

Figure 2 depicts the sequence based alignment between (A) MMP-13 and MMP-8 and (B) MMP-13 and MMP-1 used for the MMP-13 homology model.

Figure 3 is an illustration of the sulfonamide derivative of the hydroxamic inhibitor N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide (Compound A), with the corresponding proton labels.

Figure 4 lists the atomic structure coordinates for the restrained minimized mean structure of MMP-13 complexed with Compound A as derived by NMR spectroscopy. "Atom type" refers to the atom whose coordinates are being measured. "Residue" refers to the type of residue of which each measured atom is a part - i.e., amino acid, cofactor, ligand or solvent. The "x, y and z" coordinates indicate the Cartesian coordinates of each measured atom's location

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(Å). All non-protein atoms (Compound A, zinc and calcium) are listed as HETATM instead of atoms using PDB conventions.

Figure 5 lists the atomic structure coordinates for MMP-13 as derived by X-ray diffraction of a crystallized MMP-13:Compound A complex.

Figure headings are as noted above, except "Occ" indicates the occupancy factor, and "B" indicates the "B-value", which is a measure of how mobile the atom is in the atomic structure (Å<sup>2</sup>). "MOL" indicates the segment identification used to uniquely identify each molecule in the crystal.

Figure 6 is an illustration of the Compound B inhibitor, with the corresponding proton labels.

Figure 7 is a design scheme dividing 2-[Benzyl-(4-phenethyloxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide (hereinafter referred to as "Compound C") into two components corresponding to its potency component (2-[Benzyl-(4-methoxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, hereinafter referred to as "Compound D") and its selectivity component, thereby providing the basis for the design of a hybrid inhibitor with Compound B.

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Figure 8A is a design scheme showing the flow from Compound B and Compound C to the hybrid inhibitor benzofuran-2-carboxylic acid (2-{4-20 [benzyl-(2-hydroxycarbamoyl-4,6-dimethyl-phenyl)-sulfamoyl]-phenoxy}-ethyl)-amide (hereinafter referred to as "Compound E"). Figure 8B illustrates an expanded view of the NMR MMP-13:Compound B complex overlayed with the MMP-13:Compound D model, demonstrating the approach to forming the hybrid inhibitor Compound E. The MMP-13 active site is shown as a grid surface with Compound B and Compound D shown as liquorice bonds. The view is looking at the S1' pocket.

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### Detailed Description of the Invention

As used herein, the following terms and phrases shall have the meanings set forth below:

"Compound A" is N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)pyridin-3-ylmethyl-amino]-3-methyl-benzamide, as shown in Figure 3.

"Compound B" is the compound having the chemical structure shown in Figure 6. "Compound C" is 2-[Benzyl-(4-phenethyloxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, as shown in Figure 7. "Compound D" is 2-[Benzyl-(4-methoxy-benzenesulfonyl)-amino]-N-hydroxy-3,5-dimethyl-benzamide, also shown in Figure 7. "Compound E" is Benzofuran-2-carboxylic acid (2-{4-[benzyl-(2-hydroxycarbamoyl-4,6-dimethyl-phenyl)-sulfamoyl]-phenoxy}-ethyl)-amide, as shown in Figure 8A. "Compound F" is 2-(Benzyl-4-(3-phenyl-propoxy)-benzenesulfonyl]-amino)-N-hydroxy-3,5-dimethyl-

benzamide.

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Unless otherwise noted, "MMP-13" includes both human collagenase 3 as encoded by the amino acid sequence of Figure 1 (including conservative substitutions thereof), as well as "MMP-13 analogues", defined herein as proteins comprising an MMP-13 like active site as defined by the present invention, including, but not limited to, an active site characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and conserved backbone atoms of said amino acids of not more than 1.5Å; or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å. Alternatively, an MMP-13 analogue of the present invention is a protein which comprises an MMP-13 like active site characterized by a catalytic zinc, a beta strand, a Ca<sup>2+</sup> binding loop, an alpha helix and a random coil region, or, more particularly, comprising an active site characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and of amino acid residues N14, L15, T16, Y17, R18, I19, V20,

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F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, or more preferably, where said three dimensional structure further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to Figures 4 or 5, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms (N, C $\alpha$ , C, and O) of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

Unless otherwise indicated, "protein" or "molecule" shall include a protein, protein domain, polypeptide or peptide.

"Structural coordinates" are the Cartesian coordinates corresponding to an atom's spatial relationship to other atoms in a molecule or molecular complex. Structural coordinates may be obtained using x-ray crystallography techniques or NMR techniques, or may be derived using molecular replacement analysis or homology modeling. Various software programs allow for the graphical representation of a set of structural coordinates to obtain a three dimensional representation of a molecule or molecular complex. The structural coordinates of the present invention may be modified from the original sets provided in Figures 4 or 5 by mathematical manipulation, such as by inversion or integer additions or subtractions. As such, it is recognized that the structural coordinates of the present invention are relative, and are in no way specifically limited by the actual x, y, z coordinates of Figures 4 and 5. Further, it is recognized that the structural coordinates taken from Figure 5 may be from either molecule of MMP-13 catalytic fragment in the MMP-13:Compound A crystal (i.e., from A-13 or B-13).

An "agent" shall include a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug.

"Root mean square deviation" is the square root of the arithmetic mean of the squares of the deviations from the mean, and is a way of expressing deviation or variation from the structural coordinates described herein.

It will be obvious to the skilled practitioner that the numbering of the amino acid residues in the various isoforms of MMP-13 or in MMP-13 analogues covered by the present invention may be different than that set forth herein, or may contain certain conservative amino acid substitutions that yield the same three dimensional structures as those defined by Figures 4 or 5 herein. Corresponding amino acids and conservative substitutions in other isoforms or analogues are easily identified by visual inspection of the relevant amino acid sequences or by using commercially available homology software programs. "Conservative substitutions" are those amino acid substitutions which are functionally equivalent to the substituted amino acid residue, either by way of having similar polarity, steric arrangement, or by belonging to the same class as the substituted residue (e.g., hydrophobic, acidic or basic), and includes substitutions having an inconsequential effect on the three dimensional structure of MMP-13 with respect to the use of said structure for the identification and design of MMP-13 activators or inhibitors, for molecular replacement analyses and/or for homology modeling.

An "active site" refers to a region of a molecule or molecular complex that, as a result of its shape and charge potential, favorably interacts or associates with another agent (including, without limitation, a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug). As such, the active site may include both the actual site of substrate cleavage or collagenase activity, as well as certain or all binding sites or pockets adjacent to the site of substrate cleavage that nonetheless may affect MMP-13 activity upon interaction or association with an agent, either by direct interference with the site of substrate cleavage or by indirectly affecting the steric conformation or charge potential of the MMP-13 molecule. The catalytic 30 center of the MMP-13 molecule is characterized by a zinc atom chelated by H119, H123 and H129. MMP-13 binding sites or pockets located to the right of

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the catalytic zinc include S1', S2' and S3'. Binding sites or pockets to the left of the catalytic zinc include S1, S2 and S3.

The present invention relates to the three dimensional structure of human collagenase 3 (MMP-13) or an MMP-13 analogue, and more specifically, to the crystal and solution structures of MMP-13 complexed with an inhibitor, referred to herein as "Compound A", as determined using crystallography, spectroscopy and various computer modeling techniques. The three dimensional solution and crystal structures of the MMP-13:Compound A complex (as disclosed herein at Figures 4 or 5, respectively) and the uncomplexed MMP-13 catalytic fragment (which may be computationally derived from the structural coordinates of Figures 4 or 5) are useful for a number of applications, including, but not limited to, the visualization, identification and characterization of MMP-13 active sites, including the MMP-13 catalytic zinc chelated by H119, H123 and H129, as well as the various MMP-13 binding pockets adjacent to the catalytic zinc of the MMP-13 molecule. The active site structures may then be used to predict the orientation and binding affinity of a designed or selected activator or inhibitor of the MMP-13 protein. Accordingly, the invention is particularly directed to the three dimensional structure of an MMP-13 active site, including but not limited to the S1', S2', S3', S1, S2 and/or S3 binding pockets, taken separately or together 20 with the catalytic zinc of the MMP-13 molecule.

The present invention provides a solution comprising a biologically active catalytic fragment of human collagenase-3 (MMP-13) complexed with Compound A. In a particular embodiment, the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1, or conservative substitutions thereof. Preferably, the solution provided for herein comprises MMP-13 complexed with Compound A in a 1:1 molar ratio, and more preferably comprises 1 mM MMP-13 in an equimolar complex with Compound A, in a buffer comprising 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl<sub>2</sub>, 0.1mM ZnCl<sub>2</sub>, 2mM NaN<sub>3</sub>, and 10 mM deuterated DTT in either 90% H<sub>2</sub>O/10% D<sub>2</sub>O or 100% D<sub>2</sub>O, at a preferred pH of 6.5. The concentration of

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MMP-13:Compound A in the solution should be high enough to yield a good signal-to-noise ratio in the NMR spectrum, but not so high as to result in precipitation or aggregation of the protein. Further, the MMP-13 of the solution may be either <sup>15</sup>N enriched or <sup>15</sup>N, <sup>13</sup>C enriched. As exemplified below, NMR spectra from the solution of the present invention are preferably obtained at a temperature of 35°C.

The secondary structure of the catalytic fragment used in the solution of the present invention comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands, configured in the order  $\beta_{I}$ ,  $\alpha_{A}$ ,  $\beta_{II}$ ,  $\beta_{III}$ ,  $\beta_{IV}$ ,  $\beta_{V}$ ,  $\alpha_{B}$ , and  $\alpha_{C}$ . The three alpha helices correspond to residues 28-44 ( $\alpha_{A}$ ), 112-123 ( $\alpha_{B}$ ) and 153-163 ( $\alpha_{C}$ ) of Figure 1, and the five beta strands correspond to residues 83-86 ( $\beta_{I}$ ), 95-100 ( $\beta_{II}$ ), 59-66 ( $\beta_{III}$ ), 14-20 ( $\beta_{IV}$ ), and 49-53 ( $\beta_{V}$ ) of Figure 1, respectively. While the solution of the present invention comprises MMP-13 in a 1:1 molar ratio with Compound A, it is understood that one of ordinary skill in the art may devise additional solutions using alternate inhibitors or ligands in the appropriate molar concentrations, thereby preventing the auto-degradation of MMP-13 and creating a solution of sufficient stability and concentration to obtain a usable NMR spectrum.

The protein used in the solution of the present invention includes

MMP-13, as well as MMP-13 analogues, where said protein comprises an active site characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the solution coordinates of Figure 4, ± a root mean square

deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å. These residues comprise the residues most closely associated with Compound A in the MMP-13:Compound A complex, as determined from the observed NOEs between MMP-13 and Compound A (Table 1).

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Alternatively, a protein used in the solution of the present invention comprises an active site characterized by a catalytic zinc, a beta strand (comprising amino acid residues N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca<sup>2+</sup> binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and the amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figure 4, or more preferably, where said three dimensional structure further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figure 4 (incorporating an S1' pocket in the active site), or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to 20 Figure 4 (further defining a hydrophobic area at the bottom of the S1' pocket), including, in each case, conservative substitutions of said amino acids and, in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms (N, Ca, C, and O) of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). Finally, in the most preferred embodiment, the protein used in the solution of the present invention comprises the complete structural coordinates according to Figure 4,  $\pm$  a root mean square deviation from the conserved backbone atoms of said amino acids (or conservative substitutions thereof) of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å).

Also provided by the present invention is a crystallized catalytic fragment of MMP-13 complexed with Compound A. The crystal of the present invention effectively diffracts X-rays for the determination of the structural coordinates of the MMP-13:Compound A complex, and is characterized as being in orthorhombic form with space group P21212, and having unit cell parameters of a=108.3Å, b=79.8Å, and c=36.1Å. Further, the crystal complex of the present invention consists of two molecules of MMP-13:Compound A complex in the asymmetric crystal unit.

In a preferred embodiment, the MMP-13 of the crystal complex of the present invention comprises the amino acid residues of Figure 1 (or conservative substitutions thereof), and is characterized by a secondary structure comprising three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands, configured in the order  $\beta_I$ ,  $\alpha_A$ ,  $\beta_{II}$ ,  $\beta_{III}$ ,  $\beta_{IV}$ ,  $\beta_V$ ,  $\alpha_B$ , and  $\alpha_C$ . Further, the three alpha helices preferably correspond to residues 28-44 ( $\alpha_A$ ), 112-123 ( $\alpha_B$ ) and 153-163 ( $\alpha_C$ ) of Figure 1, and the five beta strands correspond to residues 83-86 ( $\beta_I$ ), 95-100 ( $\beta_{II}$ ), 59-66 ( $\beta_{III}$ ), 14-20 ( $\beta_{IV}$ ), and 49-53 ( $\beta_V$ ) of Figure 1, respectively.

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The protein used in the crystal or crystal complex of the present invention includes MMP-13, as well as MMP-13 analogues, where said protein comprises an active site characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the crystal coordinates of Figure 5,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å.

Alternatively, a protein used in the crystal or crystal complex of the present invention comprises an active site characterized by a catalytic zinc, a beta strand (comprising amino acid residues N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca<sup>2+</sup> binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions

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thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, characterized by a three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figure 5, or more preferably, where said three dimensional 10 structure further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figure 5 (incorporating an S1' pocket in the active site), or most preferably, where said three dimensional structure still further comprises the relative structural coordinates of F149 and P152 according to Figure 5 (further defining a hydrophobic area at the bottom of the S1' pocket), in each case, including conservative substitutions of the said amino acids and, in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 20 0.5Å).

Finally, in the most preferred embodiment, the protein used in the crystal of the present invention comprises the complete structural coordinates according to Figure 5,  $\pm$  a root mean square deviation from the conserved backbone atoms of said amino acids (or conservative substitutions thereof) of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å).

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Molecular modeling methods known in the art may be used to identify an active site or binding pocket of the MMP-13 molecule, MMP-13 molecular complex, or an MMP-13 analogue. Specifically, the structural coordinates provided by the present invention may be used to characterize a

three dimensional model of the MMP-13 molecule, molecular complex or MMP-13 analogue. From such a model, putative active sites may be computationally visualized, identified and characterized based on the surface structure of the molecule, surface charge, steric arrangement, the presence of reactive amino acids, regions of hydrophobicity or hydrophilicity, etc. Such putative active sites may be further refined using chemical shift perturbations of spectra generated from various and distinct MMP-13 complexes, competitive and non-competitive inhibition experiments, and/or by the generation and characterization of MMP-13 mutants to identify critical residues or characteristics of the active site.

The identification of putative active sites of a molecule or molecular complex is of great importance, as most often the biological activity of a molecule or molecular complex results from the interaction between an agent and one or more active sites of the molecule or molecular complex.

Accordingly, the active sites of a molecule or molecular complex are the best targets to use in the design or selection of activators or inhibitors that affect the activity of the molecule or molecular complex.

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The present invention is directed to an active site of MMP-13 or an MMP-13 analogue, that, as a result of its shape, reactivity, charge potential, etc., favorably interacts or associates with another agent (including, without limitation, a protein, polypeptide, peptide, nucleic acid, including DNA or RNA, molecule, compound, antibiotic or drug). As such, the active site of the present invention includes both the actual site of substrate cleavage or collagenase activity (the catalytic zinc chelated by H119, H123, and H129), as well as binding sites or pockets adjacent to the site of substrate cleavage (i.e., S1', S2', S3', S1, S2, and/or S3) that may nonetheless affect MMP-13 activity upon interaction or association with an agent, either by direct interference with the site of substrate cleavage or by indirectly affecting the steric conformation or charge potential of the MMP-13 molecule. Accordingly, the present invention is directed to an active site of the MMP-13 molecule characterized by a zinc atom chelated by H119, H123 and H129, and preferably the S1' binding pocket to the right of the catalytic zinc.

In an alternate embodiment, the active site of the present invention is characterized by the three dimensional structure comprising the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 (or conservative substitutions thereof) according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å, or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å.

Alternatively, the active site of the present invention is characterized by a catalytic zinc, a beta strand (comprising amino acid residues 10 N14, L15, T16, Y17, R18, I19, and V20 or conservative substitutions thereof), a Ca<sup>2+</sup> binding loop (comprising amino acid residues F75, D76, G77, P78, and S79 or conservative substitutions thereof), an alpha helix (comprising amino acid residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 or conservative substitutions thereof) and a random coil region (comprising amino acid residues P139, I140, and Y141 or conservative substitutions thereof), or, more particularly, is characterized by a three dimensional structure comprising the relative solution or crystal structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, respectively, or more preferably, where said three dimensional structure further comprises the relative solution or crystal structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, where said three dimensional structure still further comprises the relative solution or crystal structural coordinates of F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å

(or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

In order to use the structural coordinates generated for a crystal or solution structure of the present invention as set forth in Figures 4 and 5, respectively, it is often necessary to display the relevant coordinates as, or convert them to, a three dimensional shape or graphical representation, or to otherwise manipulate them. For example, a three dimensional representation of the structural coordinates is often used in rational drug design, molecular replacement analysis, homology modeling, and mutation analysis. This is typically accomplished using any of a wide variety of commercially available software programs capable of generating three dimensional graphical representations of molecules or portions thereof from a set of structural coordinates. Examples of said commercially available software programs include, without limitation, the following: GRID (Oxford University, Oxford, UK); MCSS (Molecular Simulations, San Diego, CA); AUTODOCK (Scripps Research Institute, La Jolla, CA); DOCK (University of California, San Francisco, CA); Flo99 (Thistlesoft, Morris Township, NJ); Ludi (Molecular Simulations, San Diego, CA); QUANTA (Molecular Simulations, San Diego, CA); Insight (Molecular Simulations, San Diego, CA); SYBYL (TRIPOS, Inc., St. Louis. MO); and LEAPFROG (TRIPOS, Inc., St. Louis, MO).

For storage, transfer and use with such programs, a machine, such as a computer, is provided for that produces a three dimensional representation of the MMP-13 molecule, a portion thereof (such as an active site or a binding site), a MMP-13 molecular complex, or an MMP-13 analogue. The machine of the present invention comprises a machine-readable data storage medium comprising a data storage material encoded with machine-readable data. Machine-readable storage media comprising data storage material include conventional computer hard drives, floppy disks, DAT tape, CD-ROM, and other magnetic, magneto-optical, optical, floptical and other media which may be adapted for use with a computer. The machine of the present invention also comprises a working memory for storing instructions for processing the

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machine-readable data, as well as a central processing unit (CPU) coupled to the working memory and to the machine-readable data storage medium for the purpose of processing the machine-readable data into the desired three dimensional representation. Finally, the machine of the present invention further comprises a display connected to the CPU so that the three dimensional representation may be visualized by the user. Accordingly, when used with a machine programmed with instructions for using said data, e.g., a computer loaded with one or more programs of the sort identified above, the machine provided for herein is capable of displaying a graphical three-dimensional representation of any of the molecules or molecular complexes, or portions of molecules of molecular complexes, described herein.

In one embodiment of the invention, the machine-readable data comprises the relative structural coordinates of the catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to Figures 4 or 5, in each case, including conservative substitutions thereof, and in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å), wherein said structural coordinates characterize an active site of MMP-13 or an MMP-13 analogue.

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In an alternate preferred embodiment, the machine-readable data comprises the structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, in each case, including conservative substitutions thereof, and in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, and most preferably, not more than 0.5Å). In an even more preferred embodiment, the machine-readable data further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126,

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L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, or most preferably, still further comprises the relative structural coordinates of F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å).

Finally, it is most preferred that the machine-readable data

comprise the relative structural coordinates of all residues constituting the

MMP-13 catalytic fragment according to Figures 4 or 5, in each case, ± a root

mean square deviation from the conserved backbone atoms of said amino acids

of not more than 1.5Å. In each case, the noted embodiments comprise

conservative substitutions of the noted residues resulting in same structural

coordinates within the stated root mean square deviation.

The structural coordinates of the present invention permit the use of various molecular design and analysis techniques in order to (i) solve the three dimensional structures of related molecules, molecular complexes or MMP-13 analogues, and (ii) to design, select, and synthesize chemical agents capable of favorably associating or interacting with an active site of an MMP-13 molecule or MMP-13 analogue, wherein said chemical agents potentially act as activators or inhibitors of MMP-13 or of an MMP-13 analogue.

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More specifically, the present invention provides a method for determining the molecular structure of a molecule or molecular complex whose structure is unknown, comprising the steps of obtaining crystals or a solution of the molecule or molecular complex whose structure is unknown, and then generating x-ray diffraction data from the crystallized molecule or molecular complex, and/or generating NMR data from the solution of the molecule or molecular complex whose structure is unknown is then compared to the x-ray diffraction data obtained from the MMP-13:Compound A crystal of the present invention.

Alternatively, the NMR data from the molecule or molecular structure whose structure is unknown is then compared with the NMR data obtained from the MMP-13:Compound A solution of the present invention. Then, molecular replacement analysis is used to conform the three dimensional structure

5 determined from the MMP-13:Compound A crystal of solution of the present invention to the x-ray diffraction data from the unknown molecule or molecular complex, or, alternatively, 2D, 3D and 4D isotope filtering, editing and triple resonance NMR techniques are used to conform the three dimensional structure determined from the MMP-13:Compound A solution of the present invention to the NMR data from the solution molecule or molecular complex.

Molecular replacement analysis uses a molecule having a known structure as a starting point to model the structure of an unknown crystalline sample. This technique is based on the principle that two molecules which have similar structures, orientations and positions will diffract x-rays similarly. A corresponding approach to molecular replacement is applicable to modeling an unknown solution structure using NMR technology. The NMR spectra and resulting analysis of the NMR data for two similar structures will be essentially identical for regions of the proteins that are structurally conserved, where the NMR analysis consists of obtaining the NMR resonance assignments and the structural constraint assignments, which may contain hydrogen bond, distance, dihedral angle, coupling constant, chemical shift and dipolar coupling constant constraints. The observed differences in the NMR spectra of the two structures will highlight the differences between the two structures and identify the corresponding differences in the structural constraints. The structure determination process for the unknown structure is then based on modifying the NMR constraints from the known structure to be consistent with the observed spectral differences between the NMR spectra.

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Accordingly, in one non-limiting embodiment of the invention, the resonance assignments for the MMP-13:Compound A complex provide the starting point for resonance assignments of MMP-13 in a new MMP-13:"unsolved agent" complex. Chemical shift perturbances in two dimensional

<sup>15</sup>N/<sup>1</sup>H spectra can be observed and compared between the MMP-13:Compound A complex and the new MMP-13:agent complex. In this way, the affected residues may be correlated with the three dimensional structure of MMP-13 as provided by the relevant residues of Figure 4. This effectively identifies the region of the MMP-13:agent complex that has incurred a structural change relative to the MMP-13:Compound A complex. The <sup>1</sup>H, <sup>15</sup>N, <sup>13</sup>C and <sup>13</sup>CO NMR resonance assignments corresponding to both the sequential backbone and sidechain amino acid assignments of MMP-13 may then be obtained and the three dimensional structure of the new MMP-13:agent complex may be generated using standard 2D, 3D and 4D triple resonance NMR techniques and NMR assignment methodology, using the MMP-13:Compound A structure, resonance assignments and structural constraints as a reference. Various computer fitting analyses of the new agent with the three dimensional model of MMP-13 may be performed in order to generate an initial three dimensional model of the new agent complexed with MMP-13, and the resulting three dimensional model may be refined using standard experimental constraints and energy minimization techniques in order to position and orient the new agent in association with the three dimensional structure of MMP-13.

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The present invention further provides that the structural

coordinates of the present invention may be used with standard homology
modeling techniques in order to determine the unknown three-dimensional
structure of a molecule or molecular complex. Homology modeling involves
constructing a model of an unknown structure using structural coordinates of
one or more related protein molecules, molecular complexes or parts thereof

(i.e., active sites). Homology modeling may be conducted by fitting common or
homologous portions of the protein whose three dimensional structure is to be
solved to the three dimensional structure of homologous structural elements in
the known molecule, specifically using the relevant (i.e., homologous) structural
coordinates provided by Figures 4 and/or 5 herein. Homology may be

determined using amino acid sequence identity, homologous secondary
structure elements, and/or homologous tertiary folds. Homology modeling can

include rebuilding part or all of a three dimensional structure with replacement of amino acids (or other components) by those of the related structure to be solved.

Accordingly, a three dimensional structure for the unknown

5 molecule or molecular complex may be generated using the three dimensional structure of the MMP-13:Compound A complex of the present invention, refined using a number of techniques well known in the art, and then used in the same fashion as the structural coordinates of the present invention, for instance, in applications involving molecular replacement analysis, homology modeling, and rational drug design.

Determination of the three dimensional structure of MMP-13 and its catalytic active site as disclosed herein is critical to the rational identification and/or design of therapeutic agents that may act as inhibitors or activators of MMP-13 enzymatic activity. Alternatively, using conventional drug assay techniques, the only way to identify such an agent is to screen thousands of test compounds, either in culture or by administration to suitable animal models in a laboratory setting, until an agent having the desired inhibitory or activating effect on a target compound is identified. Necessarily, such conventional screening methods are expensive, time consuming, and do not elucidate the method of action of the identified agent on the target compound.

However, advancing X-ray, spectroscopic and computer modeling technologies allow researchers to visualize the three dimensional structure of a targeted compound. Using such a three dimensional structure, researchers identify putative binding sites and then identify or design agents to interact with these binding sites. These agents are then screened for an activating or inhibitory effect upon the target molecule. In this manner, not only are the number of agents to be screened for the desired activity greatly reduced, but the mechanism of action on the target compound is better understood.

Accordingly, the present invention further provides a method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of using a three dimensional structure of MMP-13 as defined by the relative

structural coordinates of amino acids encoding MMP-13 to design or select a potential inhibitor or activator, and synthesizing or obtaining said potential inhibitor or activator. The inhibitor or activator may be selected by screening an appropriate database, may designed *de novo* by analyzing the steric configurations and charge potentials of an empty MMP-13 active site in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other collagenases in order to create "hybrid" activators or inhibitors. The method of the present invention is preferably used to design or select inhibitors of MMP-13 activity.

An agent that interacts or associates with an active site of MMP-13 or an MMP-13 analogue may be identified by determining an active site of MMP-13 or of the MMP-13 analogue from a three dimensional model of the MMP-13 or MMP-13 analogue, and performing computer fitting analyses to identify an agent which interacts or associates with said active site. Computer fitting analyses utilize various computer software programs that evaluate the "fit" between the putative active site and the identified agent, by (a) generating a three dimensional model of the putative active site of a molecule or molecular complex using homology modeling or the atomic structural coordinates of the active site, and (b) determining the degree of association between the putative active site and the identified agent. The degree of association may be determined computationally by any number of commercially available software programs, or may be determined experimentally using standard binding assays.

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Three dimensional models of the putative active site may be generated using any one of a number of methods known in the art, and include, but are not limited to, homology modeling as well as computer analysis of raw structural coordinate data generated using crystallographic or spectroscopy techniques. Computer programs used to generate such three dimensional models and/or perform the necessary fitting analyses include, but are not limited to: GRID (Oxford University, Oxford, UK), MCSS (Molecular Simulations, San Diego, CA), AUTODOCK (Scripps Research Institute, La Jolla,

CA), DOCK (University of California, San Francisco, CA), Flo99 (Thistlesoft, Morris Township, NJ), Ludi (Molecular Simulations, San Diego, CA), QUANTA (Molecular Simulations, San Diego, CA), Insight (Molecular Simulations, San Diego, CA), SYBYL (TRIPOS, Inc., St. Louis. MO) and LEAPFROG (TRIPOS, Inc., St. Louis, MO).

In a preferred method of the present invention, the identified active site of MMP-13 or the MMP-13 analogue comprises a catalytic zinc, a beta strand, a Ca<sup>2+</sup> binding loop, an alpha helix and a random coil region. More preferably, the identified active site comprises a catalytic zinc, a beta strand comprising residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1 (or conservative substitutions thereof), a Ca<sup>2+</sup> binding loop comprising residues F75, D76, G77, P78, and S79 according to Figure 1 (or conservative substitutions thereof), an alpha helix comprising residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1 (or conservative substitutions thereof), and a random coil region comprising residues P139, I140, and Y141 according to Figure 1 (or conservative substitutions thereof).

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More specifically, the identified active site of the present method comprises the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). In an alternate preferred embodiment, the identified active site further comprises the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case,

± a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). In yet a third preferred embodiment, the identified active site of the present method further comprises the relative structural coordinates of amino acid residues F149 and P152 according to Figures 4 or 5, in each case, including conservative substitutions of said amino acids, and in each case, ± a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å (or more preferably, not more than 1.0Å, or most preferably, not more than 0.5Å). Embodiments comprising conservative substitutions of the noted amino acids result in the same structural coordinates of the corresponding residues in Figures 4 or 5 within the stated root mean square deviation.

The effect of such an agent identified by computer fitting analyses on MMP-13 (or MMP-13 analogue) activity may be further evaluated computationally, or experimentally by contacting the identified agent with MMP-13 (or an MMP-13 analogue) and measuring the effect of the agent on the enzyme's activity. Depending upon the action of the agent on the active site of MMP-13, the agent may act either as an inhibitor or activator of MMP-13 activity. Standard enzymatic assays may be performed and the results analyzed to determine whether the agent is an inhibitor of MMP-13 activity (i.e., the agent may reduce or prevent binding affinity between MMP-13 and the relevant substrate, and thereby reduce the level or rate of MMP-13 activity compared to baseline), or an activator of MMP-13 activity (i.e., the agent may increase binding affinity between MMP-13 and the relevant substrate, and thereby increase the level or rate of MMP-13 activity compared to baseline). Further tests may be performed to evaluate the selectivity of the identified agent to MMP-13 with regard to the other metalloproteinases.

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Agents designed or selected to interact with MMP-13 must be capable of both physically and structurally associating with MMP-13 *via* various covalent and/or non-covalent molecular interactions, and of assuming a three

dimensional configuration and orientation that complements the relevant active site of the MMP-13 molecule.

Accordingly, using these criteria, the structural coordinates of the MMP-13:Compound A complex as disclosed herein, and/or structural coordinates derived therefrom using molecular replacement analysis or homology modeling, agents may be designed to increase either or both of the potency and selectivity of known inhibitors or activators, either by modifying the structure of known inhibitors or activators or by designing new agents *de novo* via computational inspection of the three dimensional configuration and electrostatic potential of an MMP-13 active site.

Accordingly, in one embodiment of the invention, the structural coordinates of Figures 4 or 5 of the present invention, or structural coordinates derived therefrom using molecular replacement or homology modeling techniques as discussed above, are used to screen a database for agents that may act as potential inhibitors or activators of MMP-13 activity (or the activity of MMP-13 analogues). Specifically, the obtained structural coordinates of the present invention are read into a software package and the three dimensional structure is analyzed graphically. A number of computational software packages may be used for the analysis of structural coordinates, including, but not limited to, Sybyl (Tripos Associates), QUANTA and XPLOR (Brunger, A.T., (1993) XPLOR Version 3.1 Manual, Yale University, New Haven, CT). Additional software programs check for the correctness of the coordinates with regard to features such as bond and atom types. If necessary, the three dimensional structure is modified and then energy minimized using the appropriate software until all of the structural parameters are at their equilibrium/optimal values. The energy minimized structure is superimposed against the original structure to make sure there are no significant deviations between the original and the energy minimized coordinates.

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The energy minimized coordinates of MMP-13 complexed with a "solved" inhibitor or activator are then analyzed and the interactions between the solved ligand and MMP-13 are identified. The final MMP-13 structure is

modified by graphically removing the solved inhibitor or activator so that only MMP-13 and a few residues of the solved agent are left for analysis of the binding site cavity. QSAR and SAR analysis and/or conformational analysis may be carried out to determine how other inhibitors or activators compare to the solved inhibitor or activator. The solved agent may be docked into the uncomplexed structure's binding site to be used as a template for data base searching, using software to create excluded volume and distance restrained queries for the searches. Structures qualifying as hits are then screened for activity using standard assays and other methods known in the art.

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Further, once the specific interaction is determined between the solved inhibitor or activator, docking studies with different inhibitors or activators allow for the generation of initial models of new inhibitors or activators in complex with MMP-13. The integrity of these new models may be evaluated a number of ways, including constrained conformational analysis using molecular dynamics methods (i.e., where both MMP-13 and the complexed activator or inhibitor are allowed to sample different three dimensional conformational states until the most favorable state is reached or found to exist between the protein and the complexed agent). The final structure as proposed by the molecular dynamics analysis is analyzed visually to make sure that the model is in accord with known experimental SAR based on measured binding affinities. Once models are obtained of the original solved agent bound to MMP-13 and computer models of other molecules bound to MMP-13, strategies are determined for designing modifications into the activators or inhibitors to improve their activity and/or enhance their selectivity.

Once an MMP-13 binding agent has been optimally selected or designed, as described above, substitutions may then be made in some of its atoms or side groups in order to improve or modify its selectivity and binding properties. Generally, initial substitutions are conservative, i.e., the replacement group will have approximately the same size, shape, hydrophobicity and charge

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original group. Such substituted chemical compounds may then be analyzed for efficiency of fit to MMP-13 by the same computer methods described in detail above.

Alternatively, the present invention provides a method for identifying a potential inhibitor or activator that is selective for one or more members of the matrix metalloproteinase family except MMP-13, comprising the steps of (i) using the three dimensional structures of MMP-13 and the desired target matrix metalloproteinase(s) as defined by the relative structural coordinates of amino acids encoding MMP-13 and the target matrix metalloproteinase(s) in order to design or select such a potential inhibitor or 10 activator, and (ii) synthesizing or obtaining said potential inhibitor or activator. In this case, the potential inhibitor or activator is designed to incorporate chemical or steric features favorable for association with an active site of the desired matrix metalloproteinase(s) and unfavorable for association with an MMP-13 active site, preferably where said active site comprises the MMP-13 S1' 15 pocket. The inhibitor or activator may be selected by screening an appropriate database, may designed de novo by analyzing the steric configurations and charge potentials of empty MMP-13/matrix metalloproteinase active sites in conjunction with the appropriate software programs, or may be designed using characteristics of known inhibitors or activators to MMP-13 or other 20 collagenases in order to create "hybrid" activators or inhibitors.

Various molecular analysis and rational drug design techniques are further disclosed in U.S. Patent Nos. 5,834,228, 5,939,528 and 5,865,116, as well as in PCT Application No. PCT/US98/16879, published as WO 99/09148, the contents of which are hereby incorporated by reference.

The present invention may be better understood by reference to the following non-limiting Examples. The following Examples are presented in order to more fully illustrate the preferred embodiments of the invention, and should in no way be construed as limiting the scope of the present invention.

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#### Example 1

<sup>1</sup>H, <sup>15</sup>N and <sup>13</sup>CO Assignments and Secondary Structure Determination of MMP-13 Complexed with Compound A

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Methods and Results: The uniform <sup>15</sup>N and <sup>13</sup>C- labeled 165 amino-acid catalytic fragment of human collagenase-3 (MMP-13) was expressed in *E. coli* strain BL21(DE3) containing the plasmid pProMMP-13 according to a published method (Freije *et al.*, <u>J. Biol. Chem.</u> 1994). MMP-13 was purified as previously described (Moy *et al.*, <u>J. Biomol.</u> 1997) with minor modifications. N-terminal amino acid sequencing was performed to confirm the protein's identity while the uniform <sup>15</sup>N and <sup>13</sup>C labeling of MMP-13 was confirmed by MALDI-TOF mass spectrometry (PerSeptive Biosystems). The sulfonamide derivative of the hydroxamic acid compound, N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide, was prepared from 2-amino-3-methyl-benzoic acid methyl ester and p-methoxybenzenesulfonyl chloride followed by alkylation with 3-picolyl chloride, hydrolysis (LiOH/THF) to afford the carboxylic acid and conversion to the hydroxamic acid (oxalyl chloride/DMF/NH2OH). Formation of the HCl salt yielded Compound A as shown in Figure 3.

The NMR samples contained 1 mM of MMP-13 determined spectrophotometrically in a equimolar complex with Compound A in a buffer containing 10 mM deuterated Tris-Base, 100 mM NaCl, 5 mM CaCl<sub>2</sub>, 0.1 mM  $\rm ZnCl_2$ , 2 mM NaN<sub>3</sub>, 10 mM deuterated DTT, in either 90% H<sub>2</sub>O/ 10% D<sub>2</sub>O or 100% D<sub>2</sub>O at pH 6.5. All NMR spectra were recorded at 35°C on a Bruker AMX-2 600 spectrometer equipped with a triple-resonance gradient probe.

Spectra were processed using the NMRPipe software package (Delaglio *et al.*, <u>J. Biomol. NMR</u> 1995) and analyzed with PIPP (Garrett *et al.*, <u>J. Magn. Reson.</u> 1991), NMRPipe and PEAK-SORT, an in-house software package. The assignments of the <sup>1</sup>H, <sup>15</sup>N, <sup>13</sup>CO, and <sup>13</sup>C resonances were based on the following experiments: CBCA(CO)NH, CBCANH, C(CO)NH, HC(CO)NH,

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HBHA(CO)NH, HNCO, HCACO, HNHA, HNCA, HCCH-COSY and HCCH-TOCSY (for reviews, see Bax et al., Methods Enzymol. 1994; and Clore & Gronenborn, Methods Enzymol. 1994). The accuracy of the MMP-13 NMR assignments was further confirmed by sequential NOEs in the <sup>15</sup>N-edited NOESY-HSQC spectra.

Prior to analysis of the MMP-13 NMR structure, the structure determination of the inhibitor-free catalytic fragment of MMP-1 has been reported (Moy et al., Biochemistry 1998; Moy et al., J. Biomol. NMR 1997) (30 simulated annealing structures deposited with Protein Data Bank, Accession No. 1AYK; restrained minimized mean structure deposited with Protein Data Bank, 10 Accession No. 2AYK). Because the MMPs are highly autocatalytic, the NMR analysis of the inhibitor-free MMP-1 was accomplished by establishing buffer conditions where the enzyme was still active but the rate of self-cleavage of the enzyme had been diminished. This was achieved by the addition of DTT which significantly diminished self-aggregation of the enzyme and by lowering the pH of the sample to 6.5, just above the pH where the enzyme was known to be inactivated because of the loss of the catalytic zinc. Under these conditions, an MMP-1 NMR sample was typically stable for 1-2 months. Unfortunately this was not the case for MMP-13, the protein rapidly degraded within a few hours which required the use of an inhibitor to assign the MMP-13 NMR resonances.

The secondary structure of the MMP-13:Compound A complex is based on characteristic NOE data involving the NH, H $\alpha$  and H $\beta$  protons from <sup>15</sup>N-edited NOESY-HSQC and <sup>13</sup>C-edited NOESY-HMQC spectra, <sup>3</sup>JHNα coupling constants from HNHA, slowly exchanging NH protons and  $^{13}$ C $\alpha$  and  $^{13}$ C $\beta$ secondary chemical shifts (for reviews, see Wishart & Sykes, Methods Enzymol. 1994; and Wuthrich, NMR of Proteins and Nucleic Acids, John Wiley & Sons, New York 1986). It was determined that the MMP-13 NMR structure in the complex is composed of three  $\alpha$ -helices corresponding to residues 28-44 (a,). 112-123 (a<sub>6</sub>) and 153-163 (a<sub>c</sub>) and a mixed parallel and anti-parallel  $\beta$ -sheet consisting of 5 strands corresponding to residues 83-86 ( $\beta_1$ ), 95-100 ( $\beta_2$ ), 59-66  $(\beta_3)$ , 14-20  $(\beta_4)$  and 49-53  $(\beta_5)$ . This is essentially identical to the secondary structure observed for other MMP structures.

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There were three distinct regions in the MMP-13:Compound A spectra where the resonance assignments are incomplete. These correspond to residues G70-Y73, P87-N91 and T144-H148. Residues T144-H148 correspond to part of the dynamic loop region previously seen in the MMP-1 structure (Moy et al., J. Biomol. NMR 1997). This suggests a similar dynamic profile for this region in the MMP-13 structure even in the presence of a high-affinity inhibitor (IC<sub>50</sub> = 33 nM). Residues P87 to N91 contain a cluster of prolines which disrupt the sequential assignment process because of the missing NH. Residues G70 to Y73 correspond to a loop region in the vicinity of the structural zinc which was readily assigned in the MMP-1 structure. The backbone and side-chain <sup>1</sup>H, <sup>15</sup>N, <sup>13</sup>C, and <sup>13</sup>CO assignments are essentially complete for the remainder of the protein.

#### Example 2

15 High Resolution Solution Structure of the Catalytic Fragment of MMP-13 Complexed with Compound A

#### Materials and Methods:

Preparation of Compound A: The sulfonamide derivative of the hydroxamic acid
compound, Compound A, was prepared according to the procedure noted in
Example 1 to yield the compound of Figure 3.

Expression of recombinant <sup>15</sup>N and <sup>13</sup>C/ <sup>15</sup>N-labeled MMP-13: A 169-residue C-terminally truncated human collagenase-3 (MMP-13) was expressed in *E. coli*.

The coding sequence of a C-terminally truncated procollagenase was amplified by PCR from the plasmid pNot3a, that contains the entire coding sequence of MMP-13 (Frieje, *et al.*, J. <u>Biol. Chem.</u> 1994). The PCR primers contained the appropriate restriction sites for ease of cloning. The construct codes for a truncated proMMP-13 with an N-terminal methionine added and a C-terminal proline at residue 169 of the native proMMP-13 sequence. The PCR amplified DNA fragment was the cloned into pET-21a (+) at the Nde I/Sal I sites,

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resulting in a recombinant plasmid designated as pProMMP-13. E. coli bacteria, BL21(DE3), containing the plasmid pProMMP-13, were grown in LB broth supplemented with 100 μg/ml ampicilin. An overnight culture was diluted 1:20 and grown at 37°C to an  $A_{600}$  of 0.6-0.8 with vigorous shaking. Isopropyl  $\beta\text{-D-}$ 5 galactoside (IPTG) was added to a final concentration of 1 mM and cultures were shaken for 3 h at 37°C. The cells were harvested by centrifugation (7000 Xg for 15 min) at 4°C, washed with PBS, and frozen at -70°C until further use.

Uniform <sup>15</sup>N and <sup>13</sup>C- labeled ProMMP-13 was obtained by growing BL21(DE3) E. coli in defined media containing 2.0 g/l [13C6, 98%+]Dglucose and 1.0 g/l [ 15N, 98%+] ammonium chloride as the sole carbon and nitrogen sources, respectively. In addition, the defined media contained M9 salts (Sambrook, et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, New York, NY 1989), trace elements, vitamins and 100 µg/ml ampicilin. Conditions for induction and growth were the same as above.

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Purification of recombinant <sup>15</sup>N and <sup>13</sup>C MMP-13: MMP-13 was purified according to Moy et al., J. Biomol. NMR 1997, with modifications as follows. Frozen cell pellets were thawed on ice. Cells were resuspended by homogenization in lysis buffer (0.1 M Tricine, pH 8.0, 10 mM EDTA, 2mM DTT, 0.5 mM PMSF). Cells were lysed by French Press (2X) followed by treatment with lysozyme (1 mg/ml; final) at room temperature for 30 min. The lysate was centrifuged at 45,000 x g for 30 minutes. The pellet was washed twice with 50 mM Tricine pH 7.5, 0.2 M NaCl<sub>2</sub>, 0.5% Triton X-100, resuspended in fresh urea buffer (20 mM Tricine, 25 pH 7.5, 8 M urea, 0.2% NaN<sub>3</sub>, 2 mM DTT) and incubated at room temperature for 1 hour. The urea solubilized protein was centrifuged at 45,000 x g for 30 min and the resultant supernatant was filtered and applied to a Hitrap-Q Sepharose (Pharmacia Biotech) anion exchange column equilibrated in 6 M urea buffer. The column was washed with urea buffer and eluted with a 0-0.25 M NaCl linear gradient. Fractions containing proMMP-13 were detected by SDS-PAGE, pooled and quickly diluted into 5-fold excess of renaturing buffer

(50 mM Tricine, pH 7.5, 0.4 M NaCl, 10 mM CaCl<sub>2</sub>, 0.1 mM ZnOAc<sub>2</sub>, 0.02% NaN<sub>3</sub>). After 2 days of dialysis against 25 volumes of renaturing buffer (with three changes), refolded proMMP-13 was concentrated to about 4-10 mg/ml in a Millipore Biomax 5 concentrator. ProMMP-13 was activated to MMP-13CAT (catalytic domain) by an overnight incubation at 37 °C in the presence of l mM p-aminophenylmercuric acetate (APMA).

The activated protein is then applied onto a Superdex-75 16/60 gel filtration column equilibrated in 2.5 mM Tris-HCl, pH 7.5, 5 mM CaCl<sub>2</sub>, 0.4 M NaCl, 2 mM DTT, 0.02% NaN<sub>3</sub> and 0.05 mM ZnOAc<sub>2</sub>. The protein is eluted and fractions containing MMP-13CAT were identified by SDS-PAGE. Peak fractions were pooled and the protein was concentrated in a Millipore Biomax concentrator to about 5 mg/ml and stored at -70 °C. N-terminal amino acid sequencing was performed to confirm the protein's identity. The uniform <sup>15</sup>N and <sup>13</sup>C labeling of MMP-13-CAT was confirmed by MALDI-TOF mass spectrometry (PerSeptive Biosystems).

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NMR Sample Preparation: The MMP-13:Compound A NMR sample contained 1mM <sup>15</sup>N-or <sup>15</sup>N/<sup>13</sup>C-labeled MMP-13 with Compound A in a 1:1 ratio. The sample was prepared by repeated buffer exchange using 20-30ml solution containing 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl<sub>2</sub>, 0.1mM ZnCl<sub>2</sub>, 2mM NaN<sub>3</sub>, 10mM deuterated DTT, and 0.2mM Compound A in either 90% H<sub>2</sub>O/10 % D<sub>2</sub>O or 100% D<sub>2</sub>O. Buffer exchange was carried out on a Millipore Ultrafree-15 Centrifugal Filter Unit. Excess Compound A was removed by additional buffer exchanges where Compound A was removed from the buffer.

NMR Data Collection: All spectra were recorded at 35°C on a Bruker AMX-2 600 spectrometer using a gradient enhanced triple-resonance <sup>1</sup>H/<sup>13</sup>C/<sup>15</sup>N probe. For spectra recorded in H<sub>2</sub>O, water suppression was achieved with the WATERGATE sequence and water-flip back pulses (Piotto, et al., <u>J. Biomol. NMR</u> 1992; Grzesiek and Bax, <u>J. Am</u>. <u>Chem</u>. <u>Soc</u>. 1993). Quadrature detection in the

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indirectly detected dimensions were recorded with States-TPPI hypercomplex phase increment (Marion, et al., J. Magn. Reson. 1989). Spectra were collected with appropriate refocusing delays to allow for 0,0 or -90,180 phase correction.

The resonance assignments and bound conformation of Compound A in the MMP-1: Compound A complex were based on the 2D <sup>12</sup>C/<sup>12</sup>C-filtered NOESY (Petros, et al., FEBS Lett. 1992; Gemmecker, et al., J. Magn. Reson. 1992), 2D <sup>12</sup>C/<sup>12</sup>C-filtered TOCSY (Petros, et al., FEBS Lett. 1992; Gemmecker, et al., J. Magn. Reson. 1992) and 12C/12C-filtered COSY experiments (Ikura and Bax, J. Magn. Reson. 1992).

The MMP-13:Compound A structure is based on the following series of spectra: HNHA (Vuister and Bax, J. Am. Chem. Soc. 1993), HNHB (Archer, et al., J. Magn. Reson. 1992), 3D long-range <sup>13</sup>C-<sup>13</sup>C correlation (Bax and Popchapsky, J. Magn. Reson. 1992), coupled CT-HCACO (Powers, et al., J. Magn. Reson. 1991; Vuister, et al., J. Am. Chem. Soc. 1992), HACAHB-COSY (Grzesiek, et al., J. Amer. Chem. Soc. 1995), 3D <sup>15</sup>N- (Mario, et al., Biochemistry 1989; Zuiderweg and Fesik, Biochemistry 1989) and <sup>13</sup>C-edited NOESY (Zuiderweg, et al., J. Magn. Reson. 1990; Ikura, et al., J. Magn. Reson. 1990), and 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY (Lee, et al., FEBS Lett. 1994). experiments. The <sup>15</sup>N-edited NOESY, <sup>13</sup>C-edited NOESY and 3D <sup>13</sup>C-edited/<sup>12</sup>Cfiltered NOESY experiments were collected with 100 msec, 120 msec and 110 msec mixing times, respectively. The acquisition parameters for each of the experiments used in determining the solution structure of MMP-13 complexed with Compound A were as reported previously (Moy, et al., Biochemistry, 1998).

Spectra were processed using the NMRPipe software package (Delaglio, et al., J. Biomol. NMR, 1995) and analyzed with PIPP (Garrett, et al., J. Magn. Reson., 1991) on a Sun Sparc Workstation. When appropriate, data processing included a solvent filter, zero-padding data to a power of two, linear predicting back one data point of indirectly acquired data to obtain zero phase 30 corrections, linear prediction of additional points for the indirectly acquired dimensions to increase resolution. Linear prediction by the means of the mirror et al., J. Mol. Biol., 1983).

image technique was used only for constant-time experiments (Zhu and Bax, <u>J</u>. <u>Magn</u>. <u>Reson</u>., 1992). In all cases data was processed with a skewed sine-bell apodization function and one zero-filling was used in all dimensions.

- Interproton Distance Restraints: The NOEs assigned from 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY and 3D <sup>15</sup>N-edited NOESY experiments were classified into strong, medium, and weak corresponding to interproton distance restraints of 1.8-2.7 Å (1.8-2.9 Å for NOEs involving NH protons), 1.8-3.3 Å (1.8-3.5 Å for NOEs involving NH protons), and 1.8-5.0 Å, respectively (Williamson, et al., J. Mol. Biol., 1985; Clore, et al., EMBO J., 1986). Upper distance limits for distances involving methyl protons and non-stereospecifically assigned methylene protons were corrected appropriately for center averaging (Wuthrich,
- Torsion Angle Restraints and Stereospecific Assignments. The β-methylene stereospecific assignments and χ<sub>1</sub> torsion angle restraints were obtained primarily from a qualitative estimate of the magnitude of <sup>3</sup>J<sub>αβ</sub> coupling constants from the HACAHB-COSY experiment (Grzesiek, et al., J. Am. Chem. Soc., 1992) and <sup>3</sup>J<sub>Nβ</sub> coupling constants from the HNHB experiment (Archer, et al., J. Magn.
   Reson., 1991). Further support for the assignments was obtained from approximate distance restraints for intraresidue NOEs involving NH, CαH, and CβH protons (Powers, et al., Biochemistry, 1993).

The φ and ψ torsion angle restraints were obtained from  ${}^3J_{NH\alpha}$  coupling constants measured from the relative intensity of Hα crosspeaks to the NH diagonal in the HNHA experiment (Vuister and Bax, J. Am. Chem. Soc. 1993), from a qualitative estimate of the magnitude of  ${}^3J_{\alpha\beta}$  coupling constants from the HACAHB-COSY experiment (Grzesiek, et al., J. Am. Chem. Soc., 1992) and from approximate distance restraints for intraresidue and sequential NOEs involving NH, CαH, and CβH protons by means of the conformational grid search program STEREOSEARCH (Nilges, et al., Biopolymers 1990), as described previously (Kraulis, et al., Biochemistry 1989).  ${}^1J_{c\alpha H\alpha}$  coupling

constants obtained from a coupled 3D CT-HCACO spectrum were used to ascertain the presence of non-glycine residues with positive f backbone torsion angles (Vuister, *et al.*, J. Am. Chem. Soc. 1992). The presence of a  $^1J_{c\alpha H\alpha}$  coupling constant greater then 130 Hz allowed for a minimum  $\varphi$  restraint of -2° to -178°.

The Ile and Leu  $\chi 2$  torsion angle restraints and the stereospecific assignments for leucine methyl groups were determined from  ${}^3J_{C\alpha C\delta}$  coupling constants obtained from the relative intensity of  $C\alpha$  and  $C\delta$  cross peaks in a 3D long-range  ${}^{13}C^{-13}C$  NMR correlation spectrum (Bax, et al., J. Am. Chem. Soc. 1992), in conjunction with the relative intensities of intraresidue NOEs (Powers, et al., Biochemistry 1993). Stereospecific assignments for valine methyl groups were determined based on the relative intensity of intraresidue NH-C $\gamma$ H and  $C\alpha$ H-C $\gamma$ H NOEs as described by Zuiderweg et al. (1985) (Zuiderweg, et al., Biopolymers 1985). The minimum ranges employed for the  $\varphi$ ,  $\psi$ , and  $\chi$  torsion angle restraints were  $\pm$  30°,  $\pm$  50°, and  $\pm$  20°, respectively (Kraulis, et al., Biochemistry 1989).

Structure Calculations: The structures were calculated using the hybrid distance geometry-dynamical simulated annealing method of Nilges *et al.* (1988) (Protein Eng.) with minor modifications (Clore, *et al.*, Biochemistry 1990) using the program XPLOR (Brunger, X-Plor Version 3.1 Manual, Yale University, New Haven, CT), adapted to incorporate pseudopotentials for  $^3J_{NH\alpha}$  coupling constants (Garrett, *et al.*, J. Magn. Reson. Ser. B 1994), secondary  $^{13}C\alpha/^{13}C\beta$  chemical shift restraints (Kuszewski, *et al.*, J. Magn. Reson. Ser B 1995) and a conformational database potential (Kuszewski, *et al.*, Protein Sci. 1996; Kuszewski, *et al.*, J. Magn. Reson. 1997). The target function that is minimized during restrained minimization and simulated annealing comprises only quadratic harmonic terms for covalent geometry,  $^3J_{NH\alpha}$  coupling constants and secondary  $^{13}C\alpha/^{13}C\beta$  chemical shift restraints, square-well quadratic potentials for the experimental distance and torsion angle restraints, and a quartic van der Waals term for non-bonded contacts. All peptide bonds were constrained to be

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planar and trans. There were no hydrogen-bonding, electrostatic, or 6-12 Lennard-Jones empirical potential energy terms in the target function.

To prevent the Zn and Ca ions from being expelled during the high-temperature simulated annealing stages of the refinement protocol, a minimal number of distance restraints between the His sidechain and Zn and between backbone atoms and Cα were included in the XPLOR distance restraint file based on the observed coordination in the X-ray structures (Lovejoy, et al., Science 1994; Lovejoy, et al., Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Borkakoti, et al., Nat. Struct. Biol. 1994).

The starting MMP-13:Compound A complex structure for the simulated-annealing protocol was obtained by manually docking Compound A into a homology model for MMP-13. The initial orientation of Compound A in the MMP-13 active site was based on the previously reported MMP-1:CGS-27023A structure (Moy, *et al.*, <u>Biochemistry</u> 1999).

Homology modeling methods were utilized to generate a three dimensional model of MMP-13. The linear amino acid sequence corresponding to the catalytic domain of MMP-13 was aligned (SYBYL) with the catalytic domains of MMP-1, MMP-7 and MMP-8 based on the availability of their x-ray crystallographic structures (Bode, et al., EMBO J 1994; Spurlino., Proteins: Struct., Funct., Genet. 1994; Betz, et al., Eur. J. Biochem. 1997; Lovejoy, et al., Nat. Struct. Biol. 1999; Borkakoti, et al., Nat. Struct. Biol. 1994; Browner, et al., Biochemistry 1995). The alignments of MMP-13 with MMP-1 and MMP-8 demonstrated the highest homology where the computed identities are 58.9% and 61.4%, respectively (Figure 2).

The X-ray structure of MMP-8 was selected to be used as the template for homology modeling the structure of MMP-13. This decision was based mainly on the sequence alignment shown in Figure 2B where no insertions (labeled "###") are found in the critical specificity loop (Labeled Underlined and Boldface). In Figure 2A, the region labeled "##" in the specificity loop shows that there is an "insertion" of 2 additional amino acid residues compared to the sequence length of MMP-1. Based on our analysis of

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the alignments, MMP-8 would allow for a more accurate modeling of the inhibitor binding pockets since no predictions have to be made within this loop region.

COMPOSER (SYBYL) was used to construct the initial homology model of MMP-13. The only insertion was a serine (labeled '\*\*' in Figure 2B) at position 32 of MMP-13. The insertion of S32 occurs within a coiled region which is at the entrance of a long alpha helix and about 17 angstroms from the S' specificity loop. The model of MMP-13 was then energy minimized utilizing a set of nested refinement procedures (Chen, *et al.*, <u>J. Biomol. Struct. Dyn.</u> 1995), but where the protein backbone heavy atoms were constrained as close as possible to their original positions.

The MMP-13:Compound A model was then subjected to a 1000 steps of CHARMM minimization with the 5 intramolecular NOE restraints and the 47 distance restraints observed between MMP-13 and Compound A where the coordinates for MMP-13 were kept fixed. This approach approximated the positioning of Compound A in the active site of MMP-13 without distorting the MMP-13 structure. The final structure was exported as a PDB file and used as the starting point for XPLOR simulated annealing protocol where all the residues in the structure were free to move. Since the initial stage of the simulated annealing protocol corresponds to high-temperature dynamics (1500 K) with a relatively weak XPLOR NOE force constant (Ries and Petrides, Biol. Chem. Hoppe-Seyler 1995), the initial MMP-13:Compound A structure does not bias the structure determination process since the structure is effectively free to explore the available conformational space. Additionally, each iteration of the simulated annealing process begins with a random trajectory for the molecular dynamics. The fact that these trajectories differ by upwards of 10 Å assures a distinct exploration of conformational space for the ensemble of MMP-13:Compound A structures determined from the simulated annealing protocol.

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#### Results and Discussion

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Compound A Resonance Assignments and Bound Conformation: The primary structure of Compound A along with the proton naming convention is shown in Figure 3. The NMR assignments for Compound A in the MMP-13 complex 5 followed established protocols using 2D <sup>12</sup>C-filtering experiments (Petros, et al., FEBS Lett. 1992; Gemmecker, et al., J. Magn. Reson. 1992; Ikura and Bax, J. Am. Chem. Soc. 1992) since the NMR sample was composed of <sup>13</sup>C/<sup>15</sup>N labeled MMP-13 and unlabeled Compound A. Thus, traditional 2D-NOESY, COSY and TOCSY spectra of Compound A in the presence of MMP-13 yielded straight-10 forward assignments for Compound A along with assignments for free Compound A (data not shown). The only notable difference in the assignments for free and bound Compound A is the observation of two distinct resonances for 2HB1/2 in the complex (4.91 ppm; 4.67 ppm). The missing resonance in the free Compound A may simply be obscured by water. Also, an observation that the protons on the p-methoxyphenyl ring are degenerate suggests rapid ring flips when complexed to MMP-13. This was also seen with CGS-27023A complexed with both MMP-1 and stromelysin (Gonnella, et al., Bioorg, Med. Chem. 1997; Moy, et al., Biochemistry 1998; Moy, et al., Biochemistry 1999).

Compound A does not adopt a preferred conformation in the absence of MMP-13 as evident by the lack of structural NOEs. Only a minimal number of intramolecular NOEs were observed for Compound A in the MMP-13 complex which were relevant to the bound conformation of Compound A (data not shown). The minimal number of structural NOEs is a result of the Compound A conformation, structure and chemical shift degeneracy. A number of the observed NOEs correspond to a sequential interaction which have no effect on the overall conformation of the inhibitor and were not used in the refinement of Compound A or the complex. The structural intramolecular NOEs observed are primarily between 1HH\* and the pyridine ring and between 2HB1/2 and both the p-methoxyphenyl and aryl ring. These NOEs are consistent with the "splayed" conformation previously observed for CGS-27023A bound to both MMP-1 and stromelysin, but the bound conformation of

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Compound A is predominately determined from the intermolecular NOEs between Compound A and MMP-13 (Table 1).

Structure Determination: The NMR structure determination methodology is an iterative procedure where the current state of the structure is used to analyze the ambiguous NOE data. In essence, the structure is used as a distance filter to sort through the ambiguous NOE list where the first structure is determined from unambiguous data. For the refinement of MMP-13, the initial structure was a homology model based on the MMP-8 X-ray structure. This was justified by the overall similarity in previously reported MMP structures and from the secondary structure assignments by NMR for MMP-13. The regular secondary structure elements of MMP-13 were identified from a qualitative analysis of sequential and inter-strand NOEs, NH exchange rates, <sup>3</sup>JHNα coupling constants (Clore, et al., Crit. Rev. Biochem. Mol. Biol. 1989) and the <sup>13</sup>Cα and <sup>13</sup>Cβ secondary chemical shifts (Spera and Bax, J. Am. Chem. Soc. 1991). The deduced secondary structure is essentially identical to the inhibitor-free MMP-1 NMR structures previously reported.

The final 30 simulated annealing structures calculated for residues 7-164 were based on 3279 experimental NMR restraints, consisting of 2561 approximate interproton distance restraints, 51 distance restraints between MMP-13 and Compound A, 88 distance restraints for 44 backbone hydrogen bonds, 391 torsion angle restraints,  $103~^3J_{NH\alpha}$  restraints 123 C $\alpha$  restraints and  $108~C\beta$  restraints. Stereospecific assignments were obtained for 81 of the 100 residues with  $\beta$ -methylene protons, for the methyl groups of 5 of the 6 Val residues, and for the methyl groups of 12 of the 13 Leu residues. In addition, 12 out of the 12 Phe residues and 7 out of the 8 Tyr residues were well defined making it possible to assign NOE restraints to only one of the pair of C $\delta$ H and C $\epsilon$ H protons and to assign a  $\epsilon$ 2 torsion angle restraint. Similarly,  $\epsilon$ 2 torsion angle restraints were assigned for the three Trp residues. The atomic rms distribution of the 30 simulated annealing structures about the mean coordinate positions for residues 7-164 is 0.43  $\pm$  0.06 Å for the backbone atoms, 0.81  $\pm$ 

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0.09 Å for all atoms, and 0.47  $\pm$  0.04 Å for all atoms excluding disordered surface side chains. The mean standard deviation for the  $\varphi$  and  $\psi$  backbone torsion angles of residues 7-164 are 6.2  $\pm$  11.3° and 7.1  $\pm$  11.8°, respectively. The high quality of the MMP-13 NMR structure is also evident by the results of PROCHECK analysis and by a calculated, large negative value for the Lennard-Jones-van der Waals energy (-695  $\pm$  11 kcal mol<sup>-1</sup>). For the PROCHECK statistics, an overall G-factor of 0.16  $\pm$  0.16, a hydrogen bond energy of 0.82  $\pm$  0.05 and only 7.8  $\pm$  1.0 bad contacts per 100 residues are consistent with a good quality structure comparable to ~1Å X-ray structure.

The high quality of the MMP-13 NMR structure is also evident by the very small deviations from idealized covalent geometry, by the absence of interproton distance and torsion angle violations greater than 0.1 Å and 1°, respectively and by the fact that most of the backbone torsion angles for non-glycine residues lie within expected regions of the Ramachandran plot (not shown). 91.5% of the residues lie within the most favored region of the Ramachandran  $\varphi$ ,  $\psi$  plot and 7.8% in the additionally allowed regions.  $^1$ JC $\alpha$ H $\alpha$  coupling constants from the coupled CT-HCACO experiment indicated that all non-glycine residues have negative  $\varphi$  torsion angles.

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The quality of the NMR data to properly define the complex is also supported by the well-defined coordinates for Compound A and the active site residues, where the atomic rms distribution is 0.47  $\pm$  0.08Å and 0.18  $\pm$  0.03 Å for the heavy atoms of Compound A and MMP-13 backbone atoms, respectively.

Description of the MMP-13:Compound A Structure: The overall fold of MMP-13 is essentially identical to previously reported MMP structures (Bode, et al., EMBO J. 1994; Gooley, et al., Nat. Struct. Biol. 1994; Lovejoy, et al., Science 1994; Lovejoy, et al., Ann. N. Y. Acad. Sci. 1994; Lovejoy, et al., Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; Stams, et al., Nat. Struct. Biol. 1994; Becker, et al., Protein Sci. 1995; Gonnella, et al., Proc. Natl. Acad. Sci. U. S. A. 1995; Van Doren, et al., Protein Sci. 1995; Botos, et al., Proc. Natl. Acad. Sci. USA 1996; Broutin, et al., Acta Crystallogr., Sect. D: Biol. Crystallogr.

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1996; Gooley, et al., J. Biomol. NMR 1996; Betz, et al., Eur. J. Biochem. 1997; Gonnella, et al., Bioorg. Med. Chem. 1997; Moy, et al., Biochemistry 1998 and Moy, et al., Biochemistry 1999). The MMP-13 NMR structure is composed of three  $\alpha$ -helices corresponding to residues 28-44 ( $\alpha_A$ ), 112-123 ( $\alpha_B$ ) and 153-163 ( $\alpha_c$ ) and a mixed parallel and anti-parallel b-sheet consisting of 5 strands corresponding to residues 83-86 ( $\beta_1$ ), 95-100 ( $\beta_2$ ), 59-66 ( $\beta_3$ ), 14-20 ( $\beta_4$ ) and 49-53 ( $\beta_{\rm s}$ ). The active site of MMP-13 is bordered by  $\beta$ -strand IV, the Ca<sup>+2</sup> binding loop, helix B and a random coil region from residues P139-Y141. The catalytic zinc is chelated by H119, H123, and H129 while the structural zinc is chelated by H69, H84 and H97. The calcium ion is chelated in a loop region consisting of residues D75 to G79. An interesting feature of the MMP active-site structure is an apparent kink in the backbone that occurs between the Ca<sup>+2</sup> binding loop and  $\beta$ -strand IV. In the case of MMP-13, this results in the NHs of both L82 and A83 facing toward the active site of the enzyme. An important feature of substrate and inhibitor binding to the MMPs are hydrogen bonding interactions with  $\beta$ -strand IV which is facilitated by this unusual kink conformation (Lovejoy, et al., Science 1994; Lovejoy, et al., Biochemistry 1994; Spurlino, et al., Proteins: Struct., Funct., Genet. 1994; and Borkakoti, et al., Nat. Struct. Biol. 1994).

The interaction of Compound A in the active site of MMP-13 was determined by 5 intramolecular NOEs for Compound A and by a total of 47 intermolecular distance restraints between MMP-13 and Compound A. The key MMP-13 residues involved in the interaction with the inhibitor correspond to three distinct MMP-13 regions: residues L81, L82 and A83 from  $\beta$ -strand IV; residues L115, V116, and H119 from  $\alpha$ -helix II; and L136, I140 and Y141 from the active site loop which comprise the S1' and S2' pockets of MMP-13. A unique feature of the MMP-13 structure is the large S1' pocket which nearly reaches the surface of the protein.

Compound A binds to the right-side of the catalytic Zn where the p-methoxyphenyl of Compound A sits in the S1' pocket of the MMP-13 active site. This positioning is evident from the observed NOEs from 3HH\*, 3HE1/2

and 3HD1/2 to L115, V116, H119, L136, and Y141. The aryl group primarily interacts with the side-chain of L81 as evident by the strong NOEs between 1HH\*, 1HE2 and 1HZ and the L81 spin-system. Finally, the pyridine ring is essentially solvent exposed but interacts with the side-chain of I140. These interactions position Compound A such that the hydroxamic acid moiety of Compound A chelates to the "right" of the catalytic zinc and the sulfonyl oxygens are in hydrogen-bonding distance to the backbone NH of L82.

It is interesting to note that the active site loop is highly dynamic in both the inhibitor-free and CGS-27023A structures based on S<sup>2</sup> orderparameters (Moy, et al., J. Biomol. NMR 1997). This region in the MMP13:Compound A structure appears to be significantly less mobile by the observation that most of the residues in this loop region were easily observable in the <sup>1</sup>H-<sup>15</sup>N HSQC spectra and readily assigned. One possible explanation for this difference is the hydrophobic interaction between the pyridine ring of
Compound A and the side-chain for Ile-140. In MMP-1, I140 is replaced by a serine which essentially eliminates this beneficial interaction.

Another unique feature of the MMP-13 NMR structure is the apparent dynamic nature of residues H69 to Y73. These residues are completely disordered due to the lack of any assignment information and the resulting absence of any constraint information presumably a result of the flexible nature of these residues. Residues H69 to Y73 occur between the Ca<sup>+2</sup> binding loop and the structural zinc where the corresponding region in the previously solved MMP-1 NMR structures is well defined. There is no apparent explanation for this change in mobility between the two NMR structures but it may contribute to the observed difference in the physical behavior of MMP-1 and MMP-13. Under identical conditions, inhibitor-free MMP-1 is stable for upwards of two months whereas inhibitor-free MMP-13 degrades immediately.

Comparison of the MMP-13:Compound A and MMP-1:CGS-27023A Structures:

The high-resolution NMR structure for the MMP-13:Compound A complex was effectively and efficiently determined by using a homology model based on the

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MMP-1 NMR structure as an initial structure to analyze ambiguous NOESY data. This result is evident of the high structural and sequence similarity between members of the MMP family and consistent with the previously observed best-fit superposition of the backbone atoms for MMP-1, stromelysin, matrilysin and neutrophil collagenase (Moy, et al., Biochemistry 1998; Moy, et al., Biochemistry 1999).

The strong similarity between the various MMP structures creates an initial difficulty in designing specific MMP inhibitors. This is exemplified by the high sequence similarity among the MMPs in the active site. Comparison of the sequence similarity between MMP-13 and MMP-1 illustrates this difficulty. There are only a few significant residue differences between the two enzymes where these modifications results in a significant change in the local environment of the active site. The R114 to V115 modification results in a conversion from a hydrophilic to a hydrophobic environment at the base of the S1' pocket between MMP-1 and MMP-13, respectively. Similarly, the N80 to L81 substitution places a bulkier hydrophobic residue in the S2' pocket for MMP-13 compared to a more hydrophilic environment for MMP-1. Similarly in the active loop region, I140 a bulky hydrophobic residue in MMP-13 replaces the smaller hydrophilic S139 residue in MMP-1. Clearly, it is feasible to incorporate substituents into a small molecule to take advantage of these spatial distinct environmental changes between MMP-1 and MMP-13. Nevertheless, when these sequence and environmental differences are averaged across the MMP family it becomes less discriminating and extremely difficult to design an inhibitor to a specific MMP subtype based strictly on the small sequence differences.

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Conversely, the most distinct structural difference between the MMPs and readily amenable to incorporating specificity in drug design is the relative size and shape of the S1' pocket. This is clearly evident by comparison of the defined S1' pockets for MMP-13 and MMP-1. The large difference in size in the S1' pockets between the MMP-13 and MMP-1 NMR structures is striking. The S1' pocket for MMP-13 nearly reaches the outer surface of the protein and

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is greater then twice the size of MMP-1. The additional size of the MMP-13 S1' pocket relative to MMP-1 is best illustrated by the filling capacity of the two inhibitors. In the MMP-1:CGS-27023A NMR structure, the p-methoxyphenyl effectively fills the available S1' pocket for MMP-1. Conversely, in the MMP-13:Compound A complex the p-methoxyphenyl only partially fills the available space within the MMP-13 S1' pocket. The size of the MMP-13 pocket is actually similar in size to stromelysin where the design of stromelysin inhibitors has taken advantage of this deeper S1' pocket by using a biphenyl substituent in another series instead of the p-methoxyphenyl in Compound A to bind into the S1' pocket (Hajduk, et al., J. Am. Chem. Soc. 1997; Olejniczak, et al., J. Am. Chem. Soc. 1997). Thus, the NMR structures for MMP-13 and MMP-1 suggest that a ready approach to designing specificity between these MMPs is to take advantage of the significantly different sized S1' pockets. The high mobility of the MMP-1 active site presents a potential caveat to this analysis of the static images of the MMP-1 and MMP-13 structures. It is probable that the MMP-1 active site is capable of accommodating a S1' substituent larger then implied from its current structure due to its increased mobility in both free and inhibited structures.

Examination of the binding mode of Compound A in the MMP13:Compound A complex suggests a conformation generally similar to CGS27023A in the MMP-1:CGS-27023A NMR structure previously reported (30 simulated annealing structures deposited with Protein Data Bank, Accession No.
4AYK; restrained minimized mean structure deposited with Protein Data Bank, Accession No. 3AYK). Compound A and CGS-27023A are structurally very similar with the only difference being the nature of the substituent binding in the S2' pocket where an aryl group in Compound A replaces the isopropyl group in CGS-27023A. The strong resemblance between the binding mode of Compound A and CGS-27023A is apparent from the nearly identical intermolecular NOE patterns observed between the inhibitors and the proteins.

The key MMP-13 residues involved in the interaction with Compound A correspond to L81, L82 and A83 from β-strand IV; residues L115, V116, and

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H119 from  $\alpha$ -helix II; and L136, I140 and Y141 from the active site loop. Similarly, the MMP-1 residues involved in the interaction with CGS-27023A correspond to residues N80, L81, A82 and H83 from  $\beta$ -strand IV; residues R114, V115, H118 and E119 from  $\alpha$ -helix II; and L135, P138, Y137, S139 and Y140 from the dynamic flexible loop.

As stated previously, there are three distinct residue changes between MMP-13 and MMP-1 in the active site. The R114 to L115 change between MMP-1 and MMP-13, respectively, has a significant impact on the environment at the base of the S1' pocket but since Compound A only partially fills the MMP-13 S1' pocket this change should not effect the binding conformation of Compound A relative to CGS-27023A. Conversely, the N80 to L81 substitution directly interacts with the inhibitors in the S2' pocket and may result in an effective change in the binding mode of the inhibitors. To complicate the analysis, the only change in the inhibitors are the substituents that bind the S2' pocket. For the MMP-1:CGS-27023A complex, the isopropyl group interacts with both the sidechains of N80 and H83 where the aryl group from Compound A only interacts with L81 in MMP-13. Additionally, CGS-27023A is in hydrogen-bonding distance to both L81 and A82, whereas Compound A appears to form a bifurcated hydrogen bond with L82. This analysis suggests that CGS-27023A binds closer to  $\beta$ -strand IV since the S2' pocket is more accessible in MMP-1 due to the absence of the bulky L81 sidechain and the presence of the aryl group in Compound A. A direct comparison of the bound conformations suggest only a subtle difference in the relative orientation of the inhibitors. The S139 to I140 difference between MMP-1 and MMP-13, respectively, appears to be related to a mobility change as opposed to a structural change. In the MMP-1:CGS-27023A structure the pyridine ring position is essentially undefined and solvent exposed this compares to the MMP-13:Compound A structure where the pyridine ring binds with the side-chain of I140. Clearly, Ile is a bulkier more hydrophobic group relative to Ser which would provide a beneficial hydrophobic interactions with the pyridine ring. The more interesting observation is the apparent decrease in mobility for the active

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loop in the MMP-13 structure which may be related the pyridine ring I140 interaction. This appears to be consistent with previously inhibited MMP X-ray structures (Spurlino, et al., Proteins: Struct., Funct., Genet. 1994) where the inhibitor may extend the formation of a  $\beta$ -sheet between b-strand IV and the active loop region which results in low B-factors in the X-ray structure. This may suggest that the mobility of the active loop region is easily removed with any positive interaction with the inhibitor.

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There are apparently some interesting differences between the mode of binding for the two inhibitors in the MMP-13: Compound A and MMP-1:CGS-27023A NMR structures. The more striking observation is the overall similarity between the two structures. Despite some significant sequence differences and a large difference in the size and shape of the S1' pocket either inhibitor structure would accurately predict the other structure. This observation seems to indicate that the major contributing factors to inhibitors binding the MMPs is the fit in the S1' pocket and the binding of the hydroxamic 15 acid to the catalytic zinc. The interaction in the S2' pocket appears to have a more subtle impact on inhibitor binding and selectivity since both Compound A and CGS-27023A are low nanomolar inhibitors of MMP-13 and MMP-1, respectively. Therefore, the high-resolution solution structure of the MMP-13:Compound A in conjunction with the previously reported MMP-1 NMR structures suggest that taking advantage of the significant differences in the size and shape of the S1' pocket is a reasonable approach for developing specific MMP inhibitors.

The studies described herein present the high-resolution solution structure of MMP-13 complexed with a sulfonamide derivative of a hydroxamic 25 acid compound (Compound A). The overall fold of MMP-13 is similar to previously reported MMPs structures. The major difference is the large S1' pocket which nearly reaches the surface of the protein. The structure was based on a total of 3279 constraints including 47 distance restraints between MMP-13 and Compound A from X-filtered NOESY experiments. The inhibitor was found 30 to bind to the "right" side of the catalytic Zn such that the p-methoxyphenyl ring

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sits in the S1' pocket, the aryl moiety interacts with L81 of βIV, the pyridine ring interacts with I140 of the active site loop, hydrogen bond interactions exist between the sulfonamide oxygens with residue L82 and the hydroxamic acid chelates the catalytic Zn. This inhibitor binds MMP-13 similarly to the MMP-1:

5 CGS-27023A complex suggesting that appropriately filling the S1' pocket may play a key role in developing selective MMP inhibitors.

Table 1. Observed NOEs Between Compound A and MMP-13

Compound A	MMP-13	NOE Class	Compound A	MMP-1	NOE Class
1HH*	L81 Hy	w	3HH*	Υ141 Ηα	.M
1HH*	L81 Hδ1#	w	3HH*	Ү141 НВ1	W
1HH*	L81 Hδ2#	М	3HH*	Υ141 Ηβ2	W
1HH*	L81 Ha	S	3HH* -	Υ141 Ηδ2	W
1HE2	L81 Hδ1#	W	3HE2	L82 Hδ1#	W
1HE2	L81 Hδ2#	М	3HE1	А83 Нβ#	W
1HZ	L81 H81#	W	3HE1	Η116 Ηα	W
1HZ	L81 Hδ2#	M	3HE1	H116 Hyl#	М
2HZ	· 1140 Ηγ2#	W	3HE2	Η116 Ηγ2#	W
2HE1 .	I140 Hδ1#	W	3HE2	1140 Ηγ2#	W
3HH*	L82 Hδ1#	W	3HE2	Υ141 Ηα	W
3HH*	L115 Hβ#	W	3HE2	Υ141 Ηβ1	W
3HH*	L115 Hy	W	3HE2	Υ141 Ηβ2	W
3HH*	L115 H81#	W	3HD2	L82 Hδ1#	W
3HH*	L115 Hδ2#	w	3HD1	· A83 Hβ#	W
3HH*	V116 Ha	w	3HD1	V116 Hy1#	w
3HH*	V116 Hy1#	W	3HD2	V116 Hγ2#	w
3HH*	V116 Hy2#	М	3HD2	1140 Ηα	w
3HH*	Η119 Ηα	w	3HD2	I140 Ηγ2#	W
3HH*	Н119 Н82	w	3HD2	Υ141 Ηα	W
3HH*	H119 Hβ1	w	3HD2	Υ141 Ηβ1	W
3HH*	Н119 Нβ2	w	3HD2	Υ141 Ηβ2	W
3HH*	L136 Hδ1#	W	3HD2	Y141 HN	w
3HH*	L136 Hδ2#	w	•		<del></del>

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## Example 3

Structure Based Design of a Novel, Potent, and Selective Inhibitor for MMP-13

The matrix metalloproteinases (MMPs) comprise a family of zinc containing enzymes that cleave a broad range of substrates including collagens, fibronectin and gelatins where the substrate preference various for individual MMPs. The design of MMP inhibitors has been initially based upon imitation of the binding interaction of natural protein substrates to MMPs where structural information of MMPs complexed with peptide substrates has been determined by x-ray crystallography and NMR spectroscopy. This structural information has provided a general description of the MMPs active site.

The active site for the MMPs is composed of a catalytic zinc chelated by three histidines where three substrate binding pockets are located to both the right (S1', S2', S3') and left (S1, S2, S3) of the catalytic zinc. The substrate binding pockets were identified by the interactions of side chains from the peptide substrate with the MMPs. The primary effort in MMP inhibitor design has focused on compounds that chelate the catalytic zinc while primarily binding in the S1' and S2' pockets. This has evolved from the observation that the structural characteristics of the S1' pocket (size, shape, amino acid composition) incurs the greatest variability between the individual MMPs and this provides an obvious approach in designing selective and specific MMP inhibitors. Nevertheless, there has also been success in utilizing the binding pockets to the left of the catalytic zinc in addition to or in combination with the right handed binding pockets in the design of inhibitors.

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The underlying challenge in designing MMP inhibitors is the reasonably high sequence and structural homology observed between the individual members of the MMP family making it intrinsically difficult to design an inhibitor that will function against a single MMP in the absence of structural information. The problem with a non-specific MMP inhibitor as a drug is the high likelihood of serious side-effects because of the large number of enzymes in the MMP family and their corresponding diversity in targets and function.

Accordingly, the detailed structural information provided herein is a critical component of an inhibitor design program targeting a particular MMP enzyme.

#### Materials and Methods:

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Synthesis of Compound D and Compound E: The sulfonamide derived from 2-amino-3,5-dimethyl-benzoic acid methyl ester and p-methoxybenzenesulfonyl chloride was N-alkylated with benzyl bromide and the ester group of the resulting intermediate was hydrolyzed (LiOH/THF) to afford the carboxylic acid. The corresponding hydroxamic acid was formed by preparation of the acid chloride (oxalyl chloride/DMF) followed by reaction with hydroxylamine. Compound E was synthesized by reaction of 2-amino-3,5-dimethyl-benzoic acid methyl ester and p-fluorobenzenesulfonyl chloride followed by N-alkylation with benzyl bromide. Hydrolysis of the methyl ester (LiOH/THF) followed by displacement of fluorine with the alkoxide of benzofuran-2-carboxylic acid (2-hydroxy-ethyl)-amide gave, after conversion to the hydroxamic acid and formation of the HCl salt as described above, Compound E.

NMR Sample Preparation: Uniformly (>95%) <sup>15</sup>N- and <sup>15</sup>N/<sup>13</sup>C-labeled human recombinant MMP-13 was expressed in *E. coli* and purified as described previously. 1mM <sup>13</sup>C/<sup>15</sup>N- and <sup>15</sup>N- MMP-13 NMR samples were prepared by concentration and buffer exchange using Millipore Ultrafree -10 centrifugal filters into a buffer containing 10mM deuterated Tris-base, 100mM NaCl, 5mM CaCl<sub>2</sub>, 0.1 mM ZnCl<sub>2</sub>, 2 mM NaN<sub>3</sub>, 10mM deuterated DTT in 90% H<sub>2</sub>O/10% D<sub>2</sub>O or 100% D2O. The 10:1 Compound B:MMP-13 samples were prepared by addition of Compound B into either a 1mM <sup>13</sup>C/<sup>15</sup>N- or <sup>15</sup>N-MMP-13 sample followed by pH readjustment. The sample to explore the potential of competitive inhibition between Compound B and Compound A was prepared by first adding 1mM of Compound A to a 1 mM <sup>15</sup>N- MMP-13 sample followed by the addition of 10mM Compound B. The initial MMP-13:Compound A sample was made by buffer exchange of <sup>15</sup>N- MMP-13 into the buffer containing 0.1 mM Compound A followed by additional buffer exchanges to remove excess

Compound A. Finally, 10mM of Compound B was added to the 1mM <sup>15</sup>N- MMP-13:Compound A sample followed by pH readjustment.

NMR Data Collection: All spectra were recorded at 35°C on a Bruker AMX-2 600
spectrometer using a gradient enhanced triple-resonance ¹H/¹³C/¹⁵N probe. For spectra recorded in H₂O, water suppression was achieved with the WATERGATE sequence and water-flip back pulses (Piotto, et al., J. Biomol. NMR 1992; Grzesiek and Bax, J. Am. Chem. Soc. 1993). Quadrature detection in the indirectly detected dimensions were recorded with States-TPPI hypercomplex
phase increment (Marion, et al., J. Magn. Reson. 1989). Spectra were collected with appropriate refocusing delays to allow for 0,0 or -90,180 phase correction.

The resonance assignments and bound conformation of Compound A in the MMP-1: Compound A complex were based on the 2D <sup>12</sup>C/<sup>12</sup>C-filtered NOESY (Petros, *et al.*, <u>FEBS Lett.</u> 1992; Gemmecker, *et al.*, <u>J. Magn. Reson.</u> 1992), 2D <sup>12</sup>C/<sup>12</sup>C-filtered TOCSY (Petros, *et al.*, <u>FEBS Lett.</u> 1992; Gemmecker, *et al.*, <u>J. Magn. Reson.</u> 1992) and <sup>12</sup>C/<sup>12</sup>C-filtered COSY experiments (Ikura and Bax, J. Am. Chem. Soc. 1992).

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The assignments of the <sup>1</sup>H, <sup>15</sup>N, and <sup>13</sup>C resonances of MMP-13 in the MMP-13:Compound B complex were based on the previous assignments for the MMP-13:Compound A complex in combination with a minimal set of experiments: 2D <sup>1</sup>H-<sup>15</sup>N HSQC, 3D <sup>15</sup>N- edited NOESY (Marion, *et al.* Biochemistry 1989; Zuiderweg and Fesik, Biochemistry 1989), CBCA(CO)NH (Grzesiek and Bax, J. Am. Chem. Soc. 1992), C(CO)NH (Grzesiek, *et al.*, J. Magn. Reson., Ser. B 1993), HNHA (Vuister and Bax, J. Am. Chem. Soc. 1993) and HNCA (Kay, *et al.*, J. Magn. Reson. 1990). The acquisition parameters for each of the experiments used in determining the solution structure of the MMP-13:Compound B complex were as reported previously (Moy, *et al.*, Biochemistry 1996).

The MMP-13:Compound B structure is based on observed NOEs from the 3D <sup>15</sup>N-edited NOESY (Marion, et al. <u>Biochemistry</u> 1989; Zuiderweg and Fesik, <u>Biochemistry</u> 1989) and 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY (Vuister

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and Bax, <u>J. Am. Chem. Soc.</u> 1993; Lee, *et al.*, <u>FEBS Lett.</u> 1994). The 3D <sup>15</sup>N-edited NOESY and 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY experiments were collected with 100 msec and 110 msec mixing times, respectively.

5 Molecular Analysis and Design: The minimized models of Compound B and Compound D complexed to MMP-13 were prepared as previously described (Chen, et al., J. Biomol. Struct. Dyn. 1995; Chen, et al., Biochemistry (in press) 1998). Using molecular dynamics methods (Sybyl v6.4 from Tripos Inc), protein regions within 5 Å from Compound B were sampled along with the inhibitor, 10 whereas everything else remained rigid during the simulations. Upon energy convergence, the last 50 frames from the final 100 picoseconds run was averaged and this averaged structure underwent a final minimization. The final protein-Compound B model appeared to have optimized possible polar and van der waals interactions. The identical procedure was applied to the complex of MMP-13 and Compound D. Since the two complexes used identical MMP-13 15 structures, the proteins were overlapped to depict the positions of the two inhibitors within the active site. Graphics analysis of the inhibitors showed that the methylene carbon of Compound B containing the 2HB1/2 protons (Figure 6) overlapped identically with the methoxy carbon from Compound D. This analysis indicated the optimal or minimal linkage length of connecting the 20 benzofuran moiety to the methoxy region of Compound D. The final design scheme is shown in Figure 8A for the hybrid inhibitor. The homology model of MMP-9 was constructed using the COMPOSER program (Tripos INC, Sybyl v.6.4)

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High-throughput Screening Analysis: Compound B was identified as an initial lead from the analysis of the MMP-13 high-throughput screen (HTS). A total of 58079 compounds were screened for their ability to inhibit MMP-13 enzymatic activity where 385 compounds were shown to have  $\geq$  40% inhibition at 10  $\mu$ g/ml dosage. Compound B was shown to exhibit weak inhibition of MMP-13 (89% at the 10  $\mu$ g/ml), but more intriguing was the observation of a complete

lack of activity against other MMPs (MMP-1, MMP-9 and TACE). The primary structure of Compound B along with the proton naming convention is shown in Figure 6.

The resulting HTS hits were further examined by cluster analysis. The hits were clustered based on structural similarities where the properties of these compounds were compared against the properties of the set of orally available drugs. The properties used to profile the HTS hits consists of: total number of non-hydrogen atoms, number of heteroatoms, number of hydrogen-bond donors and acceptors, calculated logP and molecular weight. This profile analysis provides an initial means to predict the likelihood that an HTS hit may have drug-like characteristics such as bioavailability and in-vivo stability. The profile of Compound B indicates that the compound has properties similar to orally available drugs suggesting that it would be an ideal candidate for optimization of its enzyme potency and selectivity.

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A common feature of known MMP inhibitor structures is the presence of a Zn-chelator that plays a fundamental role in its activity. In most cases Zn chelation occurs from the presence of a hydroxamic acid in the structure of the small molecule. As apparent from the structure of Compound B, the compound does not contain an obvious substituent that would chelate Zn. Thus, the unique structure of Compound B suggested a potential novel mechanism for inhibition of MMP-13 further strengthening the choice of Compound B as an initial lead candidate. Therefore, the identification of Compound B as a candidate to optimize its activity and selectivity was based on three unique observations: its intrinsic MMP-13 selectivity, its structural profile similar to known bioavailable drugs and finally its apparent novel structure.

NMR Structure of the MMP-13 - Compound B Complex: The NMR binding studies provided critical information pertaining to the mechanism of Compound B inhibition of MMP-13 and the method for designing increase potency. The major question presented when Compound B was identified from HTS was its unknown MMP-13 binding site and its method for inducing MMP-13 inhibition.

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Previous work on the NMR structure of MMP-13 complexed with Compound A and MMP-1 complexed with CGS-27023A provided the framework and methodology to analysis Compound B bound to MMP-13 (Moy, et al., Biochemistry Submitted 1999; Moy, et al., Biochemistry 1999).

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The Compound B MMP-13 binding site was initially identified from chemical shift perturbation in the <sup>1</sup>H-<sup>15</sup>N HSQC spectra. The observed perturbations were mapped onto a GRASP surface (not shown). It is apparent that the major effect of Compound B on the chemical shifts of MMP-13 occurs in the proximity of the S1' pocket suggesting that Compound B sits in this pocket. From the NMR and X-ray structures of MMP-13, it was determined that the S1' pocket for MMP-13 is very deep and linear in shape while nearly reaching the surface of the protein. In fact, a number of residues at the surface of MMP-13 near the base of the S1' pocket show significant chemical shift perturbation in the presence of Compound B. Since Compound B is a linear molecule, docking studies would place the inhibitor stretched throughout the linear S1' pocket of MMP-13. The only question remaining was whether to place the morpholine or the benzofuran moiety of Compound B at one end of the pocket, adjacent to the catalytic zinc or the opposite end, distant from the zinc atom. Property analysis of the enzymes S1' pocket depicts that the end adjacent to the zinc is relatively polar whereas the opposite end is hydrophobic. This analysis lead us to dock Compound B with the morpholine ring adjacent to the catalytic zinc atom with the benzofuran moiety siting in a hydrophobic pocket formed by L115, L136, F149 and P152 at the base of the S1' pocket. To further verify the proposed binding of Compound B in the S1' pocket of MMP-13, a simple competition experiment with Compound A was conducted. The <sup>1</sup>H-<sup>15</sup>N HSQC experiment for the MMP-13:Compound B complex was collected in the presence of Compound A. The presence of Compound A displaced all of Compound B as evident by the distinct differences in the <sup>1</sup>H-<sup>15</sup>N HSQC spectra which further suggests that both compounds bind in the S1' pocket.

The relative orientation and binding of Compound B with MMP-13 was further confirmed by the observation of intermolecular NOEs between

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Compound B and MMP-13 from the 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY experiment. The NOESY spectra was collected in the presence of a ten-fold excess of Compound B because of the weak affinity of Compound B with MMP-13. Nevertheless, a total of 16 NOEs were observed between Compound B and 5 L81, L115, V116, Y141, T142 and Y143 which support the initial positioning of Compound B in the MMP-13 S1' pocket. An expanded 2D plane from the 3D <sup>13</sup>C-edited/<sup>12</sup>C-filtered NOESY experiment (not shown) demonstrated examples of some key intermolecular NOEs between Compound B benzofuran group resonances and L115  $\delta$  and Compound B resonances proximal to the morpholine ring and L82  $\delta$ . The complex of Compound B with MMP-13 was subjected to energy refinement using the NMR results as constraints (Moy, et al., Biochemistry 1999; Chen, et al., J. Biomol. Struct. Dyn. 1995). The modeling results depict the morpholine oxygen forming a hydrogen bond with the backbone amide group of Leu-82 and the benzofuran group packs deep in the S1' pocket with the peptide bond linker portion forming hydrogen bonds 15 with protein backbone groups. The complex shows no apparent interactions between the inhibitor and the catalytic zinc justifying the ligands micromolar potency.

20 Structures of MMP-1, MMP-9 and MMP-13: The recent NMR solution structures of MMP-1 and MMP-13 were used as starting points for molecular modeling and analysis (Moy, et al., Biochemistry Submitted 1999; Moy, et al., Biochemistry 1998; Moy, et al., Biochemistry 1999). A homology model for MMP-9 was developed based on its strong homology to MMP-1 (54% identity around the catalytic domain). Based on the homology model, the catalytic site of MMP-9 is similar to the corresponding sites in MMP-1 and MMP-13. All three structures were used as starting points for analysis and synthetic design.

Comparative analysis of the MMP structures shows that residue positions 115 and 144, in addition to the length of the specificity loop, determines the size and shape of the S1' pockets. Alignment of the NMR structures for MMP-1 and MMP-13 shows that MMP-13 contains two additional

insertions in the specificity loop. The homology model of MMP-9 indicates no additional insertions so its length is identical to MMP-1.

Residue positions 115 and 144 are important in establishing the relative length of the S1' pockets for the MMPs where the larger the side chain at these positions results in a smaller S1' pocket. Since residue 115 is spatially closer to the catalytic zinc than residue 144, a larger side chain for residue 115 will have a greater impact on defining a smaller S1' pocket compared to residue 144. MMP-1 has the largest side chain at position 115, thus its S1' pocket is the smallest. MMP-9 has an Arg at position 144 resulting in its S1' pocket being longer compared to MMP-1. Conversely, MMP-13 has short side chains at both positions 115 and 144. The short side chains combined with an increased length of its specificity loop result in MMP-13 having the largest S1' pocket. To summarize, the size of the MMP S1' pockets are as follows: MMP-13 > MMP-9 > MMP-1 where this structural feature plays a critical role in the design strategy for developing a potent and specific MMP-13 inhibitor.

Design Strategy: A strategy utilizing NMR and molecular modeling was applied towards the design and synthesis of an MMP-13 selective inhibitor lead. The basic approach behind the design strategy is to optimize the affinity of the chemical lead Compound B while maintaining its inherent MMP-13 selectivity. This can be achieved by taking advantage of the distinct structural feature of MMP-13, its deep linear S1' pocket, while combining overlapping structural features of Compound B with other potent inhibitors. Compound C is an example of a potent and selective inhibitor for MMP-9 and MMP-13 (See Table 2). Based on the NMR solution structure of MMP-13 complexed with Compound A (Figure 4), structurally similar inhibitors were positioned into the active site of MMP-13.

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Figure 7 shows the critical regions of Compound C, which can be broken down into two components, Compound D which represents the zinc chelating portion of the compound that contributes to the binding potency and the toluene group (1A) which contributes to enhanced ligand selectivity against

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MMP-1. The strategy was to design a new inhibitor based on replacing the toluene group (1A) with a component of Compound B critical for binding within the extended S1' pocket of MMP-13. The overlay of the NMR solution structure for Compound B with the model for Compound D is shown in Figure 8B. The close similarity between the positioning of the two structures made it readily apparent that it would be possible to generate a hybrid of the two structures combining the potent Compound D with the selective component of Compound B (Figure 8A). These results were then used to design the proposed hybrid inhibitor Compound E. The assay data in Table 2 clearly shows that the new inhibitor, Compound E, has better potency compared to Compound C in addition to improved selectivity towards MMP-13. Thus, the combination of NMR spectroscopy with molecular modeling techniques resulted in the design of a novel, potent and selective MMP-13 inhibitor (Compound E) which has an IC50 of 17 nM for MMP-13 and showed >5800, 56 and >500 fold selectivity against MMP-1, MMP-9 and TACE, respectively. To the best of our knowledge, this represents the first example of a potent MMP-13 inhibitor that has been shown to be selective against MMP-9.

Table 2 - IC50 and Selectivity Data

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Compoun d	MIMP-1	MIMP-9	MMP-13	TACE	S-1ª	S-9*	S-TACE <sup>a</sup>		
С	750nM	46nM	75nM	470nM	10.0x	0.6x	6.3x		
D	82nM	21nM	15nM	240nM	5.5x	1.4x	16x		
Е	NA	945nM	17nM	19%	>5800x	56x	>500x		
F	1025n M	71nM	301nM	664nM	3.4x	0.2x	2.2x		
<sup>a</sup> Selectivity data presented as a ratio of the MMP or TACE IC50 with MMP-13									

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### Example 4

The X-ray crystal structure of the MMP-13:Compound A complex was determined using the following procedure:

Gene/expression system/production: The cDNA coding for human MMP-13 proenzyme had 85 residues of the PRO domain, followed by 165 residues of the catalytic domain (CAT). The gene was carried on a pET-21a expression plasmid, under the control of a bacteriophage T7 promoter. The expression host was Escherichia coli BL21(DE3), which had a chromosomal copy of T7 RNA polymerase under lac control. Cells were grown in nutrient broth, and synthesis of PRO-CAT was induced by isopropyl-β-thiogalactoside. The protein accumulated to 5-10% of total cellular protein, essentially all of which was aggregated into inclusion bodies.

For potential MAD experiments, the plasmid was transferred into a methionine auxotroph host. PRO-CAT with selenomethionine substitution was produced by induction in a defined medium, with methionine replaced by selenomethionine.

Purification and refolding of PRO-CAT: Frozen cells were disrupted
20 mechanically, and inclusion bodies were isolated by centrifugation. PRO-CAT was solubilized with urea containing dithiothreitol to disrupt any disulfide bridges. PRO-CAT was partially purified by anion-exchange chromatography, in urea, on Q Sepharose. The protein was diluted to about 400 μg/ml in a solution of sodium chloride, calcium chloride, and zinc acetate, buffered with
25 tricine-HCI. Refolding proceeded over 3-4 days, during dialysis, with multiple buffer changes. PRO-CAT was then concentrated for activation and release of CAT.

Activation of PRO-CAT: The presently-accepted view of MMPs holds that the
proenzyme form is maintained in an inactive state through the coordination of
one cysteine from the PRO domain into the active-site zinc. If this cysteine is

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displaced, the enzyme becomes active. In our protocol, aminophenyl mercuric acetate was added to the protein solution to form a mercurial adduct with the cysteine. Progress of activation was monitored by SDS polyacrylamide gel eletrophoresis. Results indicated that the CAT domain accumulated and the PRO domain was degraded to small peptides.

Purification of MMP-13 (CAT) – Size Exclusion: Following activation and PRO cleavage, MMP-13 was isolated by size-exclusion chromatography through
 SuperDex 75 in a solution of sodium chloride, calcium chloride, and zinc
 acetate, buffered with tris-HC1.

Purification of MMP-13 – Affinity: MMP-13 was further purified by affinity chromatography on an immobilized hydroxamate inhibitor. The affinity matrix was prepared by coupling an hydroxamate inhibitor to Sepharose through the amino group of the piperazine ring. MMP-13 can be absorbed to the matrix and desorbed by displacement using another inhibitor of choice.

Characterization of MMP-13: Protein preparations for crystallization trials were validated by several techniques. Routinely, SDS-PAGE showed a predominant species whose migration was consistent with a molecular weight of around 19,000. MALDITOF mass spectroscopy demonstrates a single species consistent with the expected size of 18,588 amu. (MMP-13 prepared with selenomethionine showed essentially complete replacement). N-terminal sequencing demonstrated that the protein begins with YNVF, as expected for correct cleavage between PRO and CAT. Retention volume in analytical size-exclusion chromatography was consistent with a monomeric protein: no detectable aggregation was observed. The final protein was enzymatically active on a fluorogenic peptide substrate, and degraded denatured collagen.

30 Crystallization of MMP-13 complex with Compound A: The MMP-13 protein solution was buffered with 10 mM tris-HCL buffer, pH 7.5, and 0.25 M NaCl.

The concentration of protein used for crystallization was 20.0 mg/ml. The inhibitor solution was added to a protein solution with a mole ratio (protein:inhibitor) of 1:2, and was mixed for more than 1 hour.

Crystallization conditions were screened by the hanging-drop 5 vapor diffusion method (Mcpherson, A., Methods Biochem. Anal. 1976). A successful procedure for growing crystals of this complex at room temperature was identified, and crystals were obtained. Specifically, a solution was prepared from 3 µl of protein solution and 3 µl of precipitant solution, which consisted of 26% PEG4000, 0.1 M ammonium sulfate, and 0.1 M sodium chloride. A drop of this solution was suspended on a microscope coverslip glass which had been coated with silicone to prevent drop spreading. The reservoir solutions consisted of 0.6 ml precipitant solution. Equilibration was performed at room temperature by vapor diffusion. Crystals began appearing after three days. After two weeks, these crystals stopped growing. The X-ray data which have been processed show that the MMP-13 complex was crystallized in two forms. One crystal form is C-centered orthorhombic; it belonged to space group C2221, and had a cell dimension of a=36.3 Å, b=134.4 Å, and c=134.8 Å. This crystal had high mosaicity; therefore, it would be of little use when working on the structure of the complex. The second crystal form is primitive orthorhombic, from space group P21212, with a cell constant of a=108.3 Å, b=79.8 Å, and c=36.1 Å. This crystal had low mosaicity, but it was very small in most cases.

In order to obtain a big single crystal for X-ray data collections, the seeding technique (Thaller, C., et al., J. Mol. Biol. 1981) was applied. This was accomplished by using both the microseeding and the macroseeding methods. Small seed crystals were transferred to a 20% PEG4000 precipitant solution on a depression slide. A single washed crystal was injected into a hanging-drop solution, which was composed of 3 µl of MMP-13 complex solution and 3  $\mu$ l of precipitant solution. The reservoir solutions consisted of 0.6 ml precipitant solution at pH 8.0. This procedure successfully produced bigger crystals with a maximum edge dimension of up to 0.35 x 0.1 x 0.1 mm<sup>3</sup>. These crystals diffracted X-ray at a resolution of 2.0 Å.

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X-Ray Data Collection: X-ray diffraction data from 30.0-2.0 Å resolution for the MMP-13:Compound A complex crystal (P21212 form) was collected by using an RAXIS IIc Image Plate area detector which used graphite monochromatic CuKα radiation from a Rigaku RU200 rotating anode generator (operating at 50 kV, 100 mA) at a low temperature of 100 K. The oscillation angle for each plate was 1 degree, and exposure time was 20 minutes per 'image'. The processing of X-ray diffraction data was accomplished using the HKL programs (Otwinowski, Z. and Minor, W., Methods in Enzymology 276:307-26). The R-merges for full and partial reflections were 4.0% and 6.04% respectively. 18,782 unique
reflections (81% complete at 2.0 Å resolutions) were collected.

Structure Determination and Refinement: The MMP-13 complex crystal structure has been determined by a combination of crystallographic modeling and the Molecular Replacement method using models of MMP-13 derived from the MMP-1 and MMP-8 structures. The homology between MMP-13 and MMP-8 is 56% by sequence, and at least 70% by structure. Crystals of the MMP-13 complex have two molecules in the asymmetric unit, *i.e.*, the unit is a dimer. Conventional molecular replacement was not effective for determination of this dimer structure by using a monomer model. There are two reasons for this: (1) the high symmetry of the crystal structure; and (2) the conformations and the configurations of the side chain and the main chain in flexible loops of MMP-13 and MMP-8.

Firstly, the crystal structure of the MMP-13 complex is highly symmetrical. The P21212 crystal has four symmetry operations, and there are eight molecules in a unit cell. A second crystal form, belonging to space group C222, and having eight symmetry operations in a unit cell, has been identified. In this crystal, there are 16 monomers per cell in the dimer structure, and 32 monomers per cell in the tetramer structure. Therefore, the rotation search and especial translation search become more difficult. Secondly, even though the MMP family's catalytic domain structure is highly conserved, the conformations and the configurations of the side chain and the main chain in flexible loops of

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MMP-13 and MMP-8 may not be the same. In particular, the similarity between the two structures may not be sufficient to permit the determination of the dimer structure using a monomer as the searching model.

Many attempts at a rotation and translation search were made by using the X-ray data and models of either a monomer of MMP-8 or a dimer of MMP-1. Some rotation solutions were obtained, but no final translation solution has been found by using the monomer model. Accordingly, to determine this structure, it was proposed that a dimer model be constructed first; the molecular replacement method was then applied to solve the structure.

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The key idea of this proposal was crystal packing. To construct a dimer, the orientations of each monomer were determined on the basis of a rotation search. The positions of each monomer were located on the basis of the molecular packing in unit cell. Many dimer models have been constructed and applied as the 'model' for searching the rotation and translation using program AMORE (Collaborative Computational Project, Number 4 (CCP4) (1994), Acta Cryst. D50:760-763). One dimer model was found to be correct, and finally resulted in the MMP-13 3-D crystal structure using the molecular replacement method. The MMP-13 complex structure was confirmed by observing the most important and significant fact that the positions of the two zinc ions and the two calcium ions could be identified from the difference (Fo-Fc) maps with five-sigma cut, where Fo was observed structure factor and Fc was the calculated structure factor of the dimer model without zinc and calcium atoms.

These ions were located in the exact positions where they were observed in other MMP family members. The molecule fits the (2Fo-Fc) electron densities very well, both in main chain and in side chain. The molecule fits the 2Fo-Fc electron density quite well. All of these MMP molecules are conserved in the core structure region, especially the position of the central helix and the catalytic zinc. The MMP-13 dimer structure was further confirmed by applying the molecular replacement programs XPLOR (Brünger, A.T., XPLOR Version 3.1 Manual, Yale University, New Haven CT) and MERLOT (Fitzgerald,

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P., MERLOT, version 2.4 (Nov. 10, 1991). All of them worked very well, and produced results which were in agreement with the MMP-13 structure.

Structure Refinement: The structure refinement was carried out by the program 5 XPLOR. The initial dimer model included 320 amino acid residues without zinc and calcium ions. The dimer model was refined against 2.0 Å X-ray data, collected on an RAXIS IIc area detector at a temperature of 100 K. The progress of the refinement was evaluated from the quality of the protein molecular conformations and the electron density maps, and the values of the 10 crystallographic R-factor. The initial R-factor was 52%. After rigid-body minimization, conjugated-gradient minimization, a heating stage, a slow-cooling stage in the range from 4000K to 300K, energy minimization, B-factor refinement, and positional refinement, the R-factor lowered to 0.32. Electrondensity maps with coefficients of (2Fo-Fc) and (Fo-Fc), as well as the phases, 15 were calculated. The difference map shows four zinc ions and four calcium ions in the dimer structure with five-sigma cut. Some side chain loops and a few main loops were rebuilt on the interactive graphics system. The rebuilt dimer plus the zinc and calcium ions, as the new model, was refined. The R-factor was down to 26.6%. At this stage, a model of inhibitor Compound A was 20 positioned in the active-site region based on the difference electron density.

The complex structure was refined by repeating the above steps, with the R-factor down to 20%. The water molecules were modeled as oxygen atoms. Their initial positions were located by searching the peaks in the (Fo-Fc) difference maps. These positions were then checked by calculating the distance between 'water' and the oxygen and nitrogen of the protein. Together with the protein (complex) atoms, these 'water' molecules were refined against the X-ray data. Once the temperature factor of water was higher than 50, this water was omitted. 120 water molecules near the protein were found, and five water molecules were identified in the active site of each monomer. The (2Fo-Fc) maps were used to adjust the solvent model and to aid in the placement of new solvent molecules, as well as to check and correct the whole model. The r.m.s.

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deviations of  $C\alpha$  atoms for bond angles and bond distances from ideal geometry were 1.6° and 0.012 Å. The final crystallographic R-factor was 22%, at a resolution of 2.0 Å.

All publications mentioned herein above, whether to issued

5 patents, pending applications, published articles, protein structure deposits, or
otherwise, are hereby incorporated by reference in their entirety. While the
foregoing invention has been described in some detail for purposes of clarity
and understanding, it will be appreciated by one skilled in the art from a
reading of the disclosure that various changes in form and detail can be made

10 without departing from the true scope of the invention in the appended claims.

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# What is claimed is:

- A solution comprising a biologically active catalytic 1. fragment of human collagenase-3 (MMP-13) complexed with N-Hydroxy-2-[(4methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-methyl-benzamide ("Compound A").
- 2. The solution of Claim 1, wherein the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1.
- 3. The solution of Claim 2, comprising 1 mM MMP-13 complexed with Compound A in a 1:1 molar ratio, in a buffer comprising 10mM deuterated Tris-Base, 100mM NaCl, 5mM CaCl<sub>2</sub>, 0.1mM ZnCl<sub>2</sub>, 2mM NaN<sub>3</sub>, and 10 mM deuterated DTT in either 90%  $H_2O/10\%$   $D_2O$  or 100%  $D_2O$ .
- The solution of Claim 3, wherein the MMP-13 is either 15N 4. enriched or <sup>15</sup>N, <sup>13</sup>C enriched.
- The solution of Claim 1, wherein the secondary structure of 5. the catalytic fragment of MMP-13 comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands.
- 6. The solution of Claim 5, wherein the alpha helices and beta strands are configured in the order  $\beta_{I}$ ,  $\alpha_{A}$ ,  $\beta_{II}$ ,  $\beta_{III}$ ,  $\beta_{IV}$ ,  $\beta_{V}$ ,  $\alpha_{B}$ , and  $\alpha_{C}$ .
- 7. The solution of Claim 6, wherein the three alpha helices correspond to residues 28-44 ( $\alpha_a$ ), 112-123 ( $\alpha_B$ ) and 153-163 ( $\alpha_c$ ) of Figure 1, and the five beta strands correspond to residues 83-86 ( $\beta_1$ ), 95-100 ( $\beta_{11}$ ), 59-66  $(\beta_{II})$ , 14-20  $(\beta_{IV})$ , and 49-53  $(\beta_{V})$  of Figure 1.
- A crystallized catalytic fragment of MMP-13 complexed 8. with N-Hydroxy-2-[(4-methoxy-benzenesulfonyl)-pyridin-3-ylmethyl-amino]-3-

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methyl-benzamide ("Compound A").

- 9. The crystallized complex of Claim 8, wherein the catalytic fragment of MMP-13 comprises the amino acid residues of Figure 1.
- The crystallized complex of Claim 9, characterized as being 10. in orthorhombic form with space group P21212, and having unit cell parameters of a=108.3 $\mathring{A}$ , b=79.8 $\mathring{A}$ , and c=36.1 $\mathring{A}$ .
- The crystallized complex of Claim 10, further characterized 11. as consisting of two molecules of MMP-13:Compound A complex in the asymmetric unit.
- The crystallized complex of Claim 11, wherein the 12. secondary structure of the catalytic fragment of MMP-13 comprises three alpha helices and a mixed parallel and anti-parallel beta sheet comprising five beta strands.
- 13. The crystallized complex of Claim 12, wherein the alpha helices and beta strands are configured in the order  $\beta_{I}$ ,  $\alpha_{A}$ ,  $\beta_{II}$ ,  $\beta_{IV}$ ,  $\beta_{V}$ ,  $\alpha_{B}$ , and  $\alpha_c$ .
- The crystallized complex of Claim 13, wherein the three 14. alpha helices correspond to residues 28-44 ( $\alpha_A$ ), 112-123 ( $\alpha_B$ ) and 153-163  $(\alpha_c)$  of Figure 1, and the five beta strands correspond to residues 83-86  $(\beta_t)$ , 95-100 ( $\beta_{II}$ ), 59-66 ( $\beta_{III}$ ), 14-20 ( $\beta_{IV}$ ), and 49-53 ( $\beta_{V}$ ) of Figure 1.
- 15. An active site of MMP-13, characterized by a catalytic zinc, a beta strand, a Ca<sup>2+</sup> binding loop, an alpha helix, and a random coil region.

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- 16. The active site of Claim 15, wherein the beta strand comprises residues N14, L15, T16, Y17, R18, I19, and V20 according to Figure 1, the Ca<sup>2+</sup> binding loop comprises residues F75, D76, G77, P78, and S79 according to Figure 1, the alpha helix comprises residues N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, and H123 according to Figure 1, and the random coil region comprises residues P139, I140, and Y141 according to Figure 1.
- 17. The active site of Claim 16, wherein said active site comprises the relative structural coordinates of the catalytic zinc and amino acid residues N14, L15, T16, Y17, R18, I19, V20, F75, D76, G77, P78, S79, N112, L113, F114, L115, V116, A117, A118, H119, E120, F121, G122, H123, P139, I140, and Y141 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 18. The active site of Claim 17, further comprising the relative structural coordinates of amino acid residues G80, L81, L82, A83, H84, A85, K109, G110, Y111, S124, L125, G126, L127, D128, H129, S130, K131, D132, P133, G134, A135, L136, M137, F138, T142, Y143, T144, and G145 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 19. The active site of Claim 18, further comprising the relative structural coordinates of amino acid residues F149 and P152 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å.

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- 20. An active site of MMP-13 comprising the relative structural coordinates of a catalytic zinc and amino acid residues L81, L82, L115, V116, H119, L136 and I140 according to the solution or crystal coordinates of Figures 4 or 5, respectively, in each case,  $\pm$  a root mean square deviation from the catalytic zinc and the conserved backbone atoms of said amino acids of not more than 1.5Å.
- 21. A method for identifying a potential inhibitor or activator of MMP-13, comprising the steps of:
- (a) using a three dimensional structure of MMP-13 as defined by the relative structural coordinates of amino acids encoding MMP-13 according to Figures 4 or 5,  $\pm$  a root mean square deviation from the conserved backbone atoms of said amino acids of not more than 1.5Å;
- (b) employing said three-dimensional structure to design or select a potential inhibitor or activator; and
- (c) synthesizing or obtaining said potential inhibitor or activator.
- 22. The method according to Claim 21, wherein the potential inhibitor is designed de novo.
- 23. The method according to Claim 21, wherein the potential inhibitor is designed from a known inhibitor.
- 24. The method of Claim 22, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 25. The method of Claim 23, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.

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- 26. The method according to Claim 21, wherein the step of employing the three dimensional structure to design or select the potential inhibitor comprises the steps of:
- (a) identifying chemical entities or fragments capable of associating with MMP-13; and
- (b) assembling the identified chemical entities or fragments into a single molecule to provide the structure of the potential inhibitor.
- 27. The method according to Claim 26, wherein the potential inhibitor is designed de novo.
- 28. The method according to Claim 26, wherein the potential inhibitor is designed from a known inhibitor.
- 29. The method of Claim 27, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 30. The method of Claim 28, further comprising the step of contacting the potential inhibitor with MMP-13 in the presence of a substrate to determine the ability of the potential inhibitor to inhibit MMP-13.
- 31. An inhibitor identified or designed by the method of Claim 21.
- 32. An inhibitor identified or designed by the method of Claim 26.

YNVFP	RTLKW	SKMNL	TYRIV	NYTPD
5	10	15	20	25
MTHSE	VEKAF	KKAFK	VWSDV	TPLNF
30	35	40	45	50
TRLHD	GIADI	MISFG	IKEHG	DFYPF
55	60	65	70	75
DGPSG	LLAHA	FPPGP	NYGGD	AHFDD
80	85	90	95	100
DETWT	SSSKG	YNLFL	VAAHE	FGHSL
105	110	115	120	125
GLDHS	KDPGA	LMFPI	YTYTG	KSHFM
130	135	140	145	150
LPDDD 155	VQGIQ 160	SLYG 164		

FIG. 1

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Sequence 1: MMP-13 Sequence 2: MMP-1

Identity score: 58.9 %

VGEYNVFPRTLKWSKMNLTYRIVNYTPDMTHSEVEKAFKKAFKVWSDVTPLNFTRLHDGIADIMISFGIKEHGDFYPFDG LTEGN PR WEQTHLTYRIENYTPDLPRADVDHAIEKAFQLWSNVTPLTFTKVSEGQADIMISFVRGDHRDNSPFDG

**PSGLLAHAFPPGPNYGGDAHFDDDETWTS** 

SSKGYNLF

LVAAHEFGHSLGLDHSKDPGALMF

**PIYTYTGKSHFMLPDDD**VQ

PGGNLAHAFQPGPGIGGDAHFDEDERWTNNFREYNLHRVAAHELGHSLGLS HST DIGALMY<u>PSYTFSGDYO</u>

LAODD

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GIQSLYGPGDEDPN GIQAIYGRSQ

FIG. 2A

Sequence 1: MMP-13 Sequence 2: MMP-8

Identity score: 61.4 9

VGEYNVFPRTLKWSKMNLTYRIVNYT PDMTH S EVEKAFKKAFKVWSDVTPLNFTRLHDGIADIMISFGIKEHGDFYPFDG NPKWER T NLTYRIRNYTP QLSEA EVERAI KDAFEL WSVASPLI FTRISQGEADINIAFYQRDHGDNSPFDG

PSGLLAHAFPPGPNYGGDAHFDDDETWTSSSKGYNLFLVAAHEPGHSLGLDHSKDPGALMF <u>PIYTYTGKSHFMLPDDD</u>VQ PNGILAHAFQPGQGIGGDAHFDAEETWTNTSANYNLFLVAA HEFGHSLGLAHSSDPGALMY<u>PNYAF RETSNYSLPODD</u> ID

##

GIQSLYGPGDEDPN GIQAIYG

FIG. 2B

FIG. 3

		Atom Type	Res.		x	<b>Y</b> .	Z		
ATOM	1	N	THR	7	-12.675	-13.911	-8.815	1 00	0 00
ATOM	2	HN	THR	7	-12.001	-14.254	-8.192	1.00	0.83
ATOM	3	CA	THR	· 7	-14.063	-13.649	-8.340	1.00	1.22
ATOM	4	HA	THR	ż	-14.744	-14.330	-8.830	1.00	0.63
ATOM	5	CB	THR	7	-14.132	-13.858	-6.825	1.00	0.73
ATOM	6	HB	THR	7	-13.473	-13.158	-6.335	1.00	0.66
ATOM	7	OG1	THR	7	-13.730	-15.185	-6.514	1.00	0.71
ATOM	8	HG1	THR	7	-13.721	-15.690	-7.330	1.00	1.07
MOTA	9	CG2	THR	7	-15.564	-13.628	-6.336	1.00	0.67
ATOM	10	HG21	THR	7	-15.712	-12.577	-6.139	1.00	1.14
ATOM	11	HG22	THR	7	-15.728	-14.191	-5.429	1.00	1.32
ATOM	12	HG23	THR	7	-16.261	-13.955	-7.093	1.00	1.23
ATOM	13	C	THR	7	-14.451	-12.208	-8.678	1.00	0.52
ATOM	14	0	THR	7	-15.416	-11.962	-9.374	1.00	0.65
ATOM	15	N	LEU	8	-13.704	-11.254	-8.195	1.00	0.47
MOTA	16	HN	LEU	8	-12.927	-11.473	-7.639	1.00	0.61
MOTA	17	CA	LEU	8	-14.027	-9.831	-8.495	1.00	0.42
ATOM	18	HA	LEU	8	-15.098	-9.715	-8.575	1.00	0.43
MOTA	19	CB	LEU	8	-13.495	-8.937	-7.370	1.00	0.52
ATOM	20	HB1	LEU	8	-13.721	-7.905	-7.591	1.00	0.54
ATOM	21	HB2	LEU	8	-12.424	-9.060	-7.292	1.00	0.58
MOTA	22	CG	LEU	8	-14.151	-9.331	-6.042	1.00	0.60
ATOM	23	HG	LEU	8	-13.958	-10.376	-5.844	1.00	0.60
ATOM	24	CD1	LEU	8	-13.566	-8.484	-4.910	1.00	0.74
MOTA	25	HD11	LEU	8	-13.899	-8.875	-3.960	1.00	1.22
MOTA	26	HD12	LEU	8	-13.900	-7.462	-5.016	1.00	1.26
MOTA	27	HD13	LEU	8	-12.488	-8.518	-4.956	1.00	1.31
ATOM	28	CD2	LEU	8	-15.664	-9.096	-6.117	1.00	0.61
ATOM	29	HD21	LEU	8	-15.871	-8.278	-6.791	1.00	1.13
ATOM	30	HD22	LEU	8	-16.040	-8.856	-5.134	1.00	1.18
ATOM	31	HD23	LEU	8	-16.149	-9.991	-6.478	1.00	1.26
ATOM	32	C	LEU	8	-13.374	-9.438	-9.822	1.00	0.40
ATOM	33	0	LEU	8	-12.218	-9.722	-10.064	1.00	0.45
ATOM	34	N	LYS	9	-14.109	-8.795	-10.687	1.00	0.36
MOTA	35	HN	LYS	9	-15.042	-8.581	-10.474	1.00	0.36
ATOM	36	CA	LYS	9	-13.536	-8.393	-12.002	1.00	0.37
ATOM	37	HA	LYS	9	-12.521	-8.050	-11.862	1.00	0.39
MOTA	38	CB	LYS	9	-13.539	-9.599	-12.944	1.00	0.50
MOTA	39	HB1	LYS	9	-12.851	-10.344	-12.573	1.00	0.60

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ATOM	40	HB2 1	LYS	9	-13,233	-9.286	-13.932	1.00	0.48
MOTA	41	CG I	LYS	9	-14.948	-10.193		1.00	.0.60
ATOM	42	HG1 1	LYS	9	-15.632	-9.455	-13.398	1.00	0.66
MOTA	43	HG2 1	LYS	9	-15.260	-10.482	-12.014	1.00	0.78
MOTA	44	CD 1	LYS	9 -	-14.951		-13.921	1.00	0.94
MOTA	45	HD1 1	LYS	9	-13.944	-11.794	-14.033	1.00	1.57
MOTA	46	HD2 I	LYS	9	-15.344	-11.147	-14.889	1.00	1.62
MOTA	47	CE I	LYS	9	-15.829	-12.511	-13.303	1.00	0.57
MOTA	48	HE1	LYS	9	-16.776	-12.086	-13.007	1.00	1.15
MOTA	49	HE2	LYS	9	-15.333	-12.924	-12.437	1.00	1.10
ATOM	50	NZ :	LYS	9	-16.060	-13.591	-14.304	1.00	1.61
MOTA	51	HZ1	LYS	9	-15.181	-14.127	-14.445	1.00	2.14
MOTA	52	HZ2		9		-13.168	-15.207	1.00	2.13
MOTA	53	HZ3	LYS	9	-16.802		-13.959	1.00	2.14
MOTA	54	C :	LYS	9	-14.377	-7.265	-12.605	1.00	0.32
MOTA	55	0	LYS	9	-15.493	-7.021	-12.191	1.00	0.34
MOTA	56	N	TRP	10	-13.850	-6.571	-13.577	1.00	0.31
ATOM	57	HN	TRP	10	-12.947		-13.895	1.00	0.33
MOTA	58	CA	TRP	10	-14.618	-5.456	-14.201	1.00	0.30
MOTA	59	HA	TRP	10	-15.030		-13.427	1.00	0.29
MOTA	60	CB	TRP	10	-13.684	-4.630	-15.088	1.00	0.29
MOTA	61	HB1	TRP	10	-14.264	-3.917	-15.655	1.00	0.32
MOTA	62	HB2	TRP	10	-13.157	-5.286	-15.765	1.00	0.33
MOTA	63	CG	TRP	10	-12.699	-3.901	-14.230	1.00	0.25
MOTA	64	CD1	TRP	10	-11.516		-13.812	1.00	0.30
MOTA	65		TRP	10	-11.137		-14.040	1.00	0.37
MOTA	66	CD2	TRP	10	-12.786	-2.553	-13.683	1.00	0.21
MOTA	67	NE1		10	-10.872	-3.454	-13.042	1.00	0.30
MOTA	68		TRP	10	-9.996	-3.569	-12.617	1.00	0.36
MOTA	69	CE2	TRP	10	-11.614		-12.934	1.00	0.23
MOTA	70	CE3	TRP	10	-13.758		-13.763	1.00	0.24
MOTA	71	HE3	TRP	10	-14.663	-1.706	-14.328	1.00	0.29
MOTA	72		TRP	10	-11.412	-1.075	-12.287	1.00	0.22
MOTA	73		TRP	10	-10.509	-0.903	-11.720	1.00	0.27
MOTA	74	CZ3	TRP	10	-13.558	-0.309	-13.113	1.00	0.25
MOTA	75		TRP	10	-14.310	0.463	-13.181	1.00	0.32
MOTA	76	CH2	TRP	10	-12.387	-0.078	-12.376	1.00	0.23
MOTA	77	HH2	TRP	10	-12.238	0.870	-11.879	1.00	0.26
MOTA	78	С	TRP	10	-15.755	-6.031	-15.050	1.00	0.39
MOTA	79	0	TRP	10	-15.641	-7.098	-15.620	1.00	0.48
ATOM	80	N	SER	11	-16.855		-15.132	1.00	0.43
MOTA	81	HN	SER	11	-16.927	-4.476	-14.660	1.00	0.44
MOTA	82	CA	SER	11	-18.006		-15.936	1.00	0.52
MOTA	83	HA	SER	11	-18.003	-6.915	-15.930	1.00	0.59
MOTA	84	CB	SER	11	-19.313	-5.330	-15.325	1.00	0.64
MOTA	85	HB1	SER	11	-19.120		-14.763	1.00	1.16
MOTA	86	HB2	SER	11	-19.718	-6.079	-14.666	1.00	1.20
MOTA	87	OG	SER	11	-20.246	-5.067	-16.365	1.00	1.39
MOTA	88	HG	SER	11	-19.821		-17.008	1.00	1.92
MOTA	89	C	SER	11	-17.893	-5.335	-17.379	1.00	0.47
MOTA	90	0	SER	11	-18.785			1.00	0.60
MOTA	91	N	LYS	12	-16.808	-4.692	-17.715	1.00	0.42
MOTA	92	HN	LYS	12	-16.101		-17.053	1.00	0.51
MOTA	93	CA	LYS	12	-16.646		-19.107	1.00	0.41
MOTA	94	HA	LYS	12	-17.243	-4.775	-19.781	1.00	0.47
ATOM	95	CB	LYS	12	-17.116		-19.167	1.00	0.43
MOTA	96	HB1		12	-18.168		-18.926	1.00	0.50
ATOM	97		LYS	12	-16.957		-20.163	1.00	0.46
ATOM	98	CG	LYS	12	-16.327		-18.160	1.00	0.41
MOTA	99	HG1		12	-15.275	-1.922	-18.401	1.00	0.37
MOTA	100		LYS	12	-16.484	-2.272	-17.164	1.00	0.42
MOTA	101	CD	LYS	12	-16.805	-0.430	-18.223	1.00	0.50
MOTA	102		LYS	12	-17.856		-17.981	1.00	0.56
MOTA	103		LYS	12	-16.648		-19.220	1.00	0.65
MOTA	104	CE	LYS	12	-16.018		-17.218	1.00	0.61
ATOM	105	HE1	LYS	12	-15.054		-17.636	1.00	1.15
MOTA	106		LYS	12	-15.879		-16.307	1.00	1.16
MOTA	107	NZ	LYS	12	-16.773		-16.920	1.00	1.39
ATOM	108		LYS	12	-16.498	2.018	-15.983	1.00	1.90
MOTA	109		LYS	12	-17.794	1.458	-16.927	1.00	1.87
MOTA	110			12	-16.556		-17.640	1.00	1.97
MOTA	111	Ç	LYS	12	-15.175	-4.269	-19.521	1.00	0.36
MOTA	112	0	LYS	12	-14.284	-4.250	-18.695	1.00	0.34
MOTA	113	N	MET	13	-14.917	-4.380	-20.796	1.00	0.37
atom	114	HN	MET	13	-15.652	-4.402	-21.443	1.00	0.40
ATOM	115	CA	MET	13	-13.506	-4.487	-21.269	1.00	0.38
MOTA	116	HA	MET	13	-12.910	-4.964	-20.506	1.00	0.39

ATOM	117	CB	MET	13	-13.469	-5.332 -22.543	1.00	0.46
ATOM	118	HB1	MET	13	-12.523	-5.189 -23.043		
	119	HB2	MET				1.00	0.53
MOTA				13	-14.273	-5.031 -23.199	1.00	0.42
MOTA	120	CG	MET	13	-13.632	-6.809 -22.178	1.00	0.64
ATOM	121	HG1	MET	13	-12.857	-7.097 -21.483	1.00	1.26
MOTA	122	HG2	MET	13	-13.556	-7.411 -23.071	1.00	1.37
ATOM	123	SD	MET	13				
					-15.252	-7.067 -21.414	1.00	1.22
MOTA	124	CE	MET	13	-14.663	-7.870 -19.903	1.00	0.57
ATOM	125	HEL	MET	13	-14.020	-7.189 -19.362	1.00	1.16
ATOM	126	HE2	MET	13	-14.107	-8.758 -20.158	1.00	1.09
ATOM	127		MET	13	-15.508	-8.141 -19.286		
ATOM	128	C	MET	13		-0.141 -19.260	1.00	1.20
	120				-12.936	-3.095 -21.560	1.00	0.32
ATOM	129	0	MET	13	-11.793	-2.957 -21.948	1.00	0.35
MOTA	130	N	asn	14	-13.718	-2.064 - 21.371	1.00	0.28
MOTA	131	HN	ASN	14	-14.635	-2.199 -21.052	1.00	0.29
MOTA	132	CA	ASN	14	-13.217	-0.681 -21.631		
ATOM	133	HA	ASN	14		-0.001 -21.031	1.00	0.26
					-12.359	-0.725 -22.286	1.00	0.29
MOTA	134	CB	ASN	14	-14.319	0.148 -22.297	1.00	0.30
ATOM	135	HB1	asn	14	-14.025	1.186 -22.318	1.00	0.31
MOTA	136	HB2	ASN	14	-15.235	0.043 -21.735	1.00	0.31
MOTA	137	CG	ASN	14	-14.539	-0.346 -23.729		
ATOM	138		ASN	14	13 677	-0.340 -23.729	1.00	0.37
					-13.677	-0.981 -24.304	1.00	1.16
ATOM	139		ASN	14	-15.664	-0.077 -24.334	1.00	1.05
MOTA	140	HD21		14	-16.359	0.435 -23.871	1.00	1.81
ATOM	141	HD22	ASN	14	-15.812	-0.386 <b>-</b> 25.252	1.00	1.06
ATOM	142	С	ASN	14	-12.813	-0.024 -20.309	1.00	0.22
ATOM	143	ō	ASN	14	-13.566			
ATOM	144					-0.019 -19.357	1.00	0.23
		N	LEU	15	-11.630	0.533 - 20.247	1.00	0.21
ATOM	145	HN	LEU	<b>15</b> '	-11.042	0.517 -21.031	1.00	0.24
ATOM	146	CA	LEU	15	-11.171	1.194 -18.987	1.00	0.18
ATOM	147	HA	LEU	15	-12.025	1.447 -18.379	1.00	
MOTA	148	CB	LEU	15		0.042 10.019		0.19
ATOM	149				-10.250	0.243 -18.210	1.00	0.18
			LEU	15	-9.812	0.769 -17.375	1.00	0.19
ATOM	150		LEU	15	-9.463	-0.102 -18.865	1.00	0.21
MOTA	151	CG	LEU	15	-11.046	-0.964 -17.696	1.00	0.19
ATOM	152	HG	LEU	15	-11.547	-1.442 -18.525	1.00	0.20
ATOM	153		LEU	15	-10.086	-1.961 -17.044		
ATOM	154						1.00	0.20
				15	-9.726	-1.556 -16.110	1.00	0.98
ATOM	155			15	-9.251	-2.141 -17.704	1.00	1.04
MOTA		HD13	LEU	15	-10.604	-2.890 -16.857	1.00	1.07
MOTA	157	CD2	LEU	15	-12.083	-0.513 -16.658	1.00	0.21
ATOM	158	HD21	LEU	15	-12.114	-1.228 -15.850	1.00	1.07
MOTA	159	HD22		15	-13.055	-0.456 -17.122	1.00	
MOTA	160		LEU	15	-11.814	0.457 -16.268		1.00
MOTA	161	C	LEU	15		0.437 -18.268	1.00	1.04
ATOM	162	ŏ			-10.397	2.471 -19.334	1.00	0.18
MOTA			LEU	15	-9.785	2.570 -20.380	1.00	0.20
	163	N	THR	16	-10.425	3.447 -18.460	1.00	0.18
MOTA	164	HN	THR	16	-10.929	3.338 -17.627	1.00	0.18
MOTA	165	CA	THR	16	-9.699	4.729 -18.722	1.00	0.19
ATOM	166	HA	THR	16	-9.051	4.617 -19.574		
ATOM	167	CB	THR	16		4.017 -19.574	1.00	0.20
ATOM					-10.716	5.839 -18.996	1.00	0.22
	168	HB	THR	16	-10.198	6.729 -19.315	1.00	0.24
MOTA	169		THR	16	-11.445	6.112 -17.808	1.00	0.23
MOTA	170	HG1	THR	16	-11.821	5.286 -17.495	1.00	0.98
ATOM	171	CG2	THR	16	-11.680	5.393 -20.096	1.00	0.26
ATOM	172	HG21	THR	16	-12.200	6.254 -20.489		
MOTA	173	HG22	THR	16		0.234 ~20,409	1.00	1.05
ATOM					-12.396	4.696 -19.686	1.00	1.02
	174		THR	16	<b>-11.125</b>	4.914 -20.889	1.00	1.05
ATOM	175	. <b>C</b>	THR	16	-8.864	5.100 -17.495	1.00	0.17
MOTA	176	0	THR	16	-9.157	4.687 -16.391	1.00	0.16
ATOM	177	N	TYR	17	-7.826	5.878 -17.675		
ATOM	178	HN	TYR	ī7		5.878 -17.875	1.00	0.18
ATOM	179				-7.603	6.202 -18.574	1.00	0.19
			TYR	17	-6.981	6.268 -16.507	1.00	0.17
ATOM .	180	HA	TYR	17	-7.585	6.233 ~15.615	1.00	0.17
MOTA	181	CB	TYR	17	-5.814	5.288 -16.362	1.00	0.19
MOTA	182	HB1	TYR	17	-6.194	4.278 -16.347	1.00	0.19
MOTA	183	HB2	TYR	17	-5.292	5.488 -15.438		
ATOM	184	CG	TYR	17	-4.857		1.00	0.20
ATOM	185		TYR	17		5.445 -17.520	1.00	0.22
ATOM					-5.037	4.685 -18.682	1.00	0.26
	186		TYR	17	-5.867	3.998 -18.755	1.00	0.27
ATOM	187		TYR	17	-3.782	6.336 -17.426	1.00	0.25
ATOM	188		TYR	17	-3.643	6.923 -16.530	1.00	0.26
ATOM	189	CE1	TYR	17	-4.143	4.817 -19.751	1.00	0.31
ATOM	190	HE1	TYR	17	-4.282	4.231 -20.647		
ATOM	191	CE2	TYR	ĪŻ	-2.888	6.470 -18.496	1.00	0.36
ATOM	192	HE2	TYR	17			1.00	0.30
ATOM	193	CZ	TYR		-2.059	7.158 -18.424	1.00	0.35
	733	C	111	17	-3.068	5.710 -19.658	1.00	0.32

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ATOM	194 OH TYR	17	-2.186	5.839 -20.711	1.00 0.39
ATOM	195 HH TYR	17	-1.696	5.016 -20.790	
MOTA		_			1.00 0.85
		17	-6.448	7.692 -16.690	1.00 0.19
MOTA	197 O TYR	17	-6.414	8.220 -17.784	1.00 0.21
ATOM	198 N ARG	18	-6.044	8.320 -15.616	1.00 0.19
MOTA	199 HN ARG	18	-6.089	7.874 -14.747	1.00 0.19
ATOM	200 CA ARG	18	-5,523	9.714 -15.712	
				9.714 -15.712	1.00 0.22
MOTA	201 HA ARG	18	-5.131	9.877 -16.704	1.00 0.24
MOTA	202 CB ARG	18	-6.674	10.691 -15.447	1.00 0.27
ATOM	203 HB1 ARG	18	-6.978	10.613 -14.412	1.00 0.31
MOTA	204 HB2 ARG	18	-7.507	10.442 -16.083	
					1.00 0.30
ATOM	205 CG ARG	18	-6.229	12.127 -15.733	1.00 0.35
MOTA	206 HG1 ARG	18	-5.504	12.137 -16.531	1.00 0.93
ATOM	207 HG2 ARG	18	-5.790	12.549 -14.843	1.00 0.85
MOTA	208 CD ARG	18	-7.447	12.946 -16.149	1.00 0.81
ATOM	209 HD1 ARG	18	-8.216	12.867 -15.378	
				12.007 -15.378	1.00 1.29
ATOM	210 HD2 ARG	18	-7.838	12.561 -17.068	1.00 1.63
MOTA	211 NE ARG	18	-7.030	14.362 ~16.406	1.00 1.52
MOTA	212 HE ARG	18	-7.071	14.711 -17.318	1.00 2.11
MOTA	213 CZ ARG	18	-6.561	15.119 -15.456	1.00 2.24
MOTA	214 NH1 ARG	· 18			
_			-6.119	16.314 -15.736	1.00 3.18
ATOM	215 HH11 ARG	18	-6.142	16.647 -16.679	1.00 3.48
MOTA	216 HH12 ARG	18	-5.760	16.898 -15.009	1.00 3.84
MOTA	217 NH2 ARG	18	-6.564	14.700 -14.220	1.00 2.63
ATOM	218 HH21 ARG	18	-6.928	13.795 -14.000	1.00 2.44
MOTA	219 HH22 ARG	18			
			-6.205	15.285 -13.493	1.00 3.49
MOTA	220 C ARG	18	-4.413	9.931 -14.676	1.00 0.21
MOTA	221 O ARG	18	-4.550	9.576 -13.522	1.00 0.23
ATOM	222 N ILE	19	-3.314	10.514 -15.079	1.00 0.21
MOTA	223 HN ILE	19	-3.223		
_					1.00 0.22
ATOM	224 CA ILE	19	-2.196	10.755 -14.118	1.00 0.23
MOTA	225 HA ILE	19	-2.200	9.985 -13.360	1.00 0.25
MOTA	226 CB ILE	19	-0.864	10.721 -14:875	1.00 0.25
MOTA	227 HB ILE	19	-0.862	11.491 -15.633	1.00 0.25
MOTA	228 CG1 ILE	19		0 341 15 531	
			-0.702	9.341 -15.531	1.00 0.29
ATOM	229 HG11 ILE	19	-1.607	9.092 -16.065	1.00 0.82
ATOM	230 HG12 ILE	19	-0.525	8.601 -14.765	1.00 0.97
ATOM	231 CG2 ILE	19	0.291	10.962 -13.893	1.00 0.29
ATOM	232 HG21 ILE	19	1.231		
					1.00 1.08
ATOM		19	0.272	10.206 -13.123	1.00 1.09
ATOM	234 HG23 ILE	19	0.187	11.937 -13.440	1.00 1.00
ATOM	235 CD1 ILE	19	0.477	9.345 -16.512	1.00 0.93
ATOM	236 HD11 ILE	19	1.402	9.216 -15.970	1.00 1.59
ATOM	237 HD12 ILE	19	0.501		
ATOM					1.00 1.50
		19	0.360	8.533 -17.214	1.00 1.55
MOTA	239 C ILE	19	-2.381	12.126 -13.454	1.00 0.23
ATOM	240 O ILE	19	-2.355	13.150 -14.108	1.00 0.23
MOTA	241 N VAL	20	-2.563	12.152 -12.161	1.00 0.25
ATOM	242 HN VAL	20	-2.578	11.314 -11.653	1.00 0.27
MOTA	243 CA VAL	20	-2.746	13 454 11 454	
ATOM				13.454 -11.454	1.00 0.27
	244 HA VAL	20	-3.496	14.035 -11.970	1.00 0.27
MOTA	245 CB VAL	20	-3.202	13.205 -10.015	1.00 0.31
ATOM	246 HB VAL	20	-2.522	12.517 -9.534	1.00 0.32
ATOM	247 CG1 VAL	20	-3.216	14.529 -9.247	1.00 0.33
ATOM	248 HG11 VAL	20	-3.607	15.310 -9.883	
ATOM	249 HG12 VAL		-3.007		
ATOM		20	-2.211	14.782 -8.944	1.00 1.08
	250 HG13 VAL	20	-3.842	14.432 -8.372	1.00 1.10
MOTA	251 CG2 VAL	20	-4.612	12.611 -10.028	1.00 0.33
ATOM	252 HG21 VAL	20	-5.296	13.317 -10.476	1.00 1.05
ATOM	253 HG22 VAL	20	-4.924	12.401 -9.016	1.00 1.03
MOTA	254 HG23 VAL	20	-4.612		
MOTA				11.697 -10.602	1.00 1.11
		20	-1.424	14.231 -11.451	1.00 0.27
ATOM	256 O VAL	20	-1.403	15.435 -11.611	1.00 0.26
MOTA	257 n asn	21	-0.321	13.555 -11.259	1.00 0.28
MOTA	258 HN ASN	21	-0.357	12.585 -11.124	
ATOM	259 CA ASN	21			
ATOM			0.992	14.265 -11.235	1.00 0.29
	260 HA ASN	21	0.973	15.076 -11.949	1.00 0.26
ATOM	261 CB ASN	21	1.235	14.829 -9.834	1.00 0.33
ATOM	262 HB1 ASN	21	0.544	15.637 -9.646	1.00 0.33
ATOM	263 HB2 ASN	21	2.249		
MOTA	264 CG ASN	21			1.00 0.35
			1.022	13.727 -8.795	1.00 0.40
MOTA	265 OD1 ASN	21	0.459	12.694 -9.097	1.00 1.01
MOTA	266 ND2 ASN	21	1.445	13.908 -7.574	1.00 0.88
MOTA	267 HD21 ASN	21	1.895	14.743 -7.330	1.00 1.50
MOTA	268 HD22 ASN	21	1.312	13.208 -6.901	1.00 0.88
MOTA	269 C ASN	21	2.116	13.291 -11.606	
MOTA	270 O ASN	21			1.00 0.34
	2.0 O nan	~1	1.929	12.090 -11.619	1.00 0.37

MOTA	271	N	TYR	22	3.274	13.810 -1	1.933	1.00	0.38
ATOM	272	HN	TYR	22	3.387		1.932	1.00	0.38
ATOM	273	CA	TYR	22	4.417				
							2.340	1.00	0.46
ATOM	274	HA	TYR	22	4.067	11.929 -1	2.509	1.00	0.45
MOTA	275	CB	TYR	22	5.028	13.481 -1	3.630	1.00	0.49
ATOM	276	HB1	TYR	22	5.845	12.846 -1	3 938	1.00	0.56
	277		TYR						
MOTA		HB2		22	5.397		3.457	1.00	0.53
ATOM	278	CG	TYR	22	3.981	13.513 -1	4.714	1.00	0.43
MOTA	279	CD1	TYR	22	3.684	12.352 -1	5.436	1.00	0.38
ATOM	280	HD1	TYR	22	4.199				
						11.430 -1		1.00	0.39
MOTA	281	CD2	TYR	22	3.313	14.708 -1		1.00	0.46
MOTA	282	HD2	TYR	22	3.543	15.603 -1	4.445	1.00	0.51
ATOM	283	CE1	TYR	22	2.718	12.386 -1	6.447	1.00	0.36
MOTA	284	HE1							0.30
			TYR	22	2.490	11.491 -1		1.00	0.36
MOTA	285	CE2	TYR	22	2.345	14.742 -1	6.013	1.00	0.44
MOTA	286	HE2	TYR	22	1.828		6.235	1.00	0.49
MOTA	287	CZ	TYR	22	2.048		6.735		0.39
						13.301 -1	0.733	1.00	
MOTA	288	OH	TYŖ	22	1.095	13.615 -1	.7.733	1.00	0.43
MOTA	289	HH	TYŔ	22	1.173	14.457 -1	8.187	1.00	0.92
ATOM	290	C	TYR	22	5.499		1.258	1.00	0.56
ATOM	291	ŏ				10.323 -1	1.230		
			TYR	22	6.554	12.378 -1	1.470	1.00	1.38
ATOM	292	N	THR	23	5.240	13.544 -1	.0.130	1.00	0.47
ATOM	293	HN	THR	23	4.372		0.023	1.00	1.08
ATOM	294	CA	THR	23	6.237		9.004		
								1.00	0.46
ATOM	295	HA	THR	23	5.848		8.304	1.00	0.48
MOTA	296	CB	THR	23	6.361	12.265 -	8.273	1.00	0.62
MOTA	297	HB	THR .	23	5.383	11.969 -	7.921	1.00	0.68
MOTA	298		THR			12.303 -	7.321		
				23	7.223		7.156	1.00	0.86
MOTA	299	HG1	THR	23	7.941	11.788 -	7.244	1.00	1.28
MOTA	300	CG2	THR	23	6.916		9.181	1.00	0.59
ATOM	301	HG21	THR	23	7.753	11.533 -			
						11.555 -	9.748	1.00	1.08
ATOM	302	HG22	THR	23	6.141		-9.850	1.00	1.16
ATOM	303	HG23	THR	23	7.245	10.332 -	-8.570	1.00	1.22
MOTA	304	С	THR	23	7.623		9.523	1.00	0.40
ATOM	305	ŏ	THR						
				23	8.077		.0.565	1.00	0.45
MOTA	306	N	PRQ	24	8.302	15.016 -	-8.823	1.00	0.42
MOTA	307	CA	PRO	24	9.625		9.311	1.00	0.42
MOTA	308	HA	PRO	24	9.534				
							10.307	1.00	0.46
ATOM	309	CB	PRO	24	9.924		-8.335	1.00	0.50
ATOM	310	HB1	PRO	24	9.743	17.605 -	8.815	1.00	0.57
MOTA	311	HB2	PRO	24	10.955		-8.014	1.00	0.49
ATOM	312	CG	PRO			10.330			
				24	8.995		-7.129	1.00	0.66
MOTA	313		PRO	24	8.613	17.475 -	-6.842	1.00	0.84
MOTA	314	HG2	PRO	24	9.537		-6.303	1.00	0.76
ATOM	315	CD	PRO	24	7.832				
ATOM	316						-7.529	1.00	0.56
			PRO	24	7.675	14.826 -	-6.786	1.00	0.62
MOTA	317	HDl	PRO	24	6.940	16.183 -	-7.680	1.00	0.61
MOTA	· 318	С	PRO	24	10.743		-9.253	1.00	0.40
MOTA	319	ŏ	PRO	24	11.835				
ATOM							-9.737	1.00	0.40
	320	N	ASP	25	10.490		-8.662	1.00	0.44
ATOM	321	HM	ASP	25	9.608	13.172 -	-8.270	1.00	0.48
ATOM	322	CA	ASP	25	11.554		-8.577		
1001								1.00	0.48
ATOM	323	HA	ASP	25	12.393	12.695 -	-8.025	1.00	0.51
MOTA	324	CB	ASP	25	11.016		-7.847	1.00	0.57
MOTA	325	HB1	ASP	25	11.719	10.249 -	-7.945	1.00	0.61
MOTA	326	HB2	ASP	25	10.068		-8.276	1.00	0.56
MOTA	327	CG	ASP	25	10.827				
							-6.364	1.00	0.67
ATOM	328		ASP	25	10.079	10.689 -	-5.709	1.00	1.23
MOTA	329	OD2	ASP	25	11.437		-5.908	1.00	1.34
ATOM	330	C	ASP	25	12.025	11.916 -	-9.985	1.00	
ATOM	331	ŏ	ASP		12 150				0.45
				25	13.179	11.597 -		1.00	0.55
MOTA	332	N	MET	26	11.146	11.948 -1	LO.955	1.00	0.40
MOTA	333	HN	MET	26	10.220	12.209 -	10.767	1.00	0.41
ATOM	334	CA	MET	26	11,553	11 500	12 240		
						11.590 -		1.00	0.42
ATOM	335	HA	MET	26	12.624	11.686 -		1.00	0.49
MOTA	336	CB	MET	26	11.144	10.149 -		1.00	0.53
MOTA	337	HB1	MET	26	11.282	9.954 -		1.00	
MOTA	338		MET						0.55
				26	10.105	10.006 -		1.00	0.51
MOTA	339	CG	MET	26	12.011	9.186 -:		1.00	0.71
MOTA	340	HG1	MET	26 -	11.783	9.288 -	10 706	1.00	0.73
MOTA	341		MET	26		0.410	10.770		
					13.053	9.419 -		1.00	0.77
MOTA	342	SD	MET	26	11.683	7.485 -	12.380	1.00	0.89
MOTA	343	CE	MET	26	10.000	7.330 -	11.728	1.00	.0.59
MOTA	344		MET	26	9.292	7.456 -	12 524	1.00	
ATOM	345	HE2							1.25
				26	9.825	8.084 -	10.979	1.00	1.23
ATOM	346	HE3		26	9.877	6.352 -	11.285	1.00	1.23
MOTA	347	С	MET	26	10.872	12.530 -	3 3//	1.00	0.34
								2.00	0.34

ATOM	348	0	MET	26	9.897	13.184 -13.031	1.00	0.32
ATOM	349	N	THR	27	11.385	12.604 -14.544	1.00	0.33
ATOM	350							
		HN	THR	27	12.174	12.070 -14.773	1.00	0.38
ATOM	351	CA	THR	27	10.775	13.504 -15.562	1.00	0.32
ATOM	352	HA	THR	27	10.618	14.483 -15.133	1.00	0.35
MOTA	353	CB	THR	27	11.711			
							1.00	0.39
ATOM	354	HB	THR	27	11.295	14.308 -17.484	1.00	0.42
MOTA	355	OG1	THR	27	11.852	12.338 -17.371	1.00	0.37
ATOM	356	HG1	THR	27	12.765	12.242 -17.653		
							1.00	0.94
MOTA	357	CG2	THR	27	13.080	14.121 -16.313	1.00	0.51
ATOM	358	HG21	THR	27	13.602	14.553 -17.154	1.00	1.14
ATOM	359	HG22	THR	27	13.655	13.297 -15.918		
ATOM							1.00	1.11
		HG23	THR	27	12.951	14.871 -15.546	1.00	1.12
MOTA	361	С	THR	27	9.436	12.921 -16.013	1.00	0.27
ATOM	362	0	THR	27	9.177	11.743 -15.864	1.00	0.24
MOTA	363	Ň						0.24
			HIS	28	8.580	13.740 -16.554	1.00	0.32
MOTA	364	HN	HIS	28	8.807	14.688 -16.657	1.00	0.37
ATOM	365	CA	HIS	28	7.253	13.241 -17.004	1.00	0.34
MOTA	366	HA	HIS	28	6.715			
							1.00	0.36
MOTA	367	CB	HIS	28	6.457	14.403 -17.601	1.00	0.46
MOTA	368	HB1	HIS	28	5.428	14.104 -17.736	1.00	0.71
MOTA	369	HB2	HIS	28	6.880	14.676 -18.557	1.00	0.88
ATOM	370	CG	HIS	28	6.516			
ATOM						15.583 -16.669	1.00	0.73
	371		HIS	28 .	6.056	16.838 -17.036	1.00	1.66
ATOM	372		HIS	28	5.659	17.080 -17.898	1.00	2.30
ATOM	373	CD2	HIS	28	6.987	15.716 -15.387	1.00	1.33
MOTA	374	HD3	HIS	28		14 000 14 700		
		1102	ura		7.423	14.922 -14.798	1.00	2.01
MOTA	375	CEI	HIS	28	6.258	17.664 -15.993	1.00	. 1.95
ATOM	376	HE1	HIS	28	5.993	18.711 -15.990	1.00	2.70
ATOM	377		HIS	28	6.823	17.031 -14.962		
ATOM							1.00	1.71
	378	C	HIS	28	7.436	12.156 -18.069	1.00	0.30
ATOM	379	0	HIS	28	6.737	11.164 -18.082	1.00	0.30
ATOM	380	N	SER	29	8.362	12.338 -18.970	1.00	0.31
ATOM	381	HN	SER	29	8.912	13.149 -18.952		
							1.00	0.34
MOTA	382	CA	SER	29	8.567	11.319 -20.039	1.00	0.32
MOTA	383	HA	SER	29	7.660	11.217 -20.615	1.00	0.35
ATOM	384	CB	SER	. 29	9.699	11.775 -20.959	1.00	0.38
ATOM	385	HB1	SER	29	9.973			
						10.963 -21.621	1.00	0.39
ATOM	386	HB2	SER	29	10.555	12.056 -20.368	1.00	0.37
ATOM	387	OG	SER	29	9.265	12.896 -21.717	1.00	0.45
ATOM	388	HG	SER	29	9.157	12.614 -22.628	1.00	0.96
ATOM	389	C	SER	29		0.054 10.404		
					8.931	9.964 -19.424	1.00	0.26
MOTA	390	0	SER	29	8.479	8.930 -19.876	1.00	0.26
MOTA	391	N	GLU	30	9.747	9.954 -18.405	1.00	0.24
ATOM	392	HN	GLU	30	10.107	10.796 -18.056		
MOTA	393	CA	GLU				1.00	0.25
				30	10.137	8.657 -17.779	1.00	0.22
MOTA	394	HA	GLU	30	10.484	7.978 -18.542	1.00	0.25
MOTA	395	CB	GLU	30	11.260	8.899 -16.769	1.00	0.23
ATOM	396	HB1		30	11.424	8.002 -16.191		0.24
ATOM	397	HB2					1.00	
				30	10.980	9.707 -16.108	1.00	0.22
ATOM	398	CG	GLŲ	30	12.547	9.268 -17.510	1.00	0.29
ATOM	399	HG1	GLU	30	12.386	10.165 -18.086	1.00	0.67
ATOM	400	HG2	GLU	30	12.826	8.460 -18.171	1.00	
MOTA	401	CD	GLU			0.400 -16.171		0.68
				30	13.666	9.509 -16.495	1.00	0.84
ATOM	402		GLU	30	13.436	9.266 -15.321	1.00	1.49
MOTA	403	OE2	GLU	30	14.731	9:936 -16.908	1.00	1.59
ATOM	404	C	GLU	30	8.935	8.046 -17.051	1.00	0.17
ATOM	405	ŏ	GLU	30	8.715	E.040 44 000		
MOTA						6.849 -17.082	1.00	0.19
	406	N	VAL	31	8.163	8.861 -16.387	1.00	0.16
MOTA	407	HN	VAL	31	8.366	9,819 -16.371	1.00	0.17
atom	408	CA	VAL	31	6.983	8.341 -15.640	1.00	0.16
ATOM	409	HA	VAL	31		7 507 14 000		0.10
ATOM	410				7.292	7.527 -14.999	1.00	0.17
		CB	VAL	31	6.402	9:464 -14.782	1.00	0.20
MOTA	411	HB	VAL	31	6.261	10,344 -15.392	1.00	0.22
MOTA	412	CG1	VAL	31	5.058	9.021 -14.208	1.00	0.23
MOTA		HG11		31		0 000 -3 0C=		
					5.135	8.000 -13.867	1.00	0.97
MOTA		HG12		31	4.298	9:090 -14.973	1.00	1.07
MOTA		HG13		31	4.793	9.659 -13.378	1.00	1.07
ATOM	416	CG2	VAL	31	7.364	9.785 -13.636		
MOTA		HG21		31		3.103 -T3.030	1.00	0.24
					7.528	8.897 -13.045	1.00	1.05
ATOM	418			31	6.936	10.557 -13.013	1.00	1.03
ATOM	419		VAL	31	8.304	10.129 -14.040	1.00	0.99
ATOM	420	С	VAL	31	5.911	7.844 -16.617		
MOTA	421	ŏ	VAL	31			1.00	0.16
					5.293	6.817 -16.406	1.00	0.17
MOTA	422	N	GLU	32	5.672	8.571 -17.677	1.00	0.18
MOTA	423	HN	GLU	32	6.172	9.401 -17.824	1.00	0.19
MOTA	424	CA	GLU	32	4.626	8.146 -18.652	1.00	0.13
					4.020	D.140 -18.632	4 . 000	

ATOM	425	HA GLU	32	. 3.673	8.092 -18.147	1.00	0 24
ATOM	426	CB GLU		4.533			0.24
ATOM	427	HB1 GLU				1.00	0.27
				3.922	8.772 -20.582	1.00	0.31
MOTA	428	HB2 GLU		5.524	9.379 -20.164	1.00	0.28
MOTA	429	CG GLU		3.904	10.463 -19.262	1.00	0.29
MOTA	430	. HG1 GLU	32	4.456	10.812 -18.405	1.00	0.48
MOTA	431	HG2 GLU		2.879	10.272 -18.977	1.00	0.52
MOTA	432	CD GLU		3.937			
	433					1.00	0.70
MOTA		OE1 GLU		4.969	12.161 -20.513	1.00	1.37
MOTA	434	OE2 GLU		2.929	11.696 -21.026	1.00	1.45
MOTA	435	C GL		4.962	6.773 -19.235	1.00	0.20
MOTA	436	O GLU	32	4.126	5.893 -19.280	1.00	0.20
MOTA	437	N LYS		6.168	6.575 -19.689	1.00	0.20
ATOM	438	HN LYS		6.835	7.293 -19.654		
ATOM	439	CA LY			7.293 -19.034	1.00	0.21
MOTA				6.518	5.249 -20.269	1.00	0.21
	440	HA LYS		5.825	5.029 -21.068	1.00	0.24
MOTA	441	CB LYS		7.940	5.281 -20.843	1.00	0.26
atom	442	HB1 LYS		7.987	6.024 -21.624	1.00	0.31
ATOM	443	HB2 LYS	33	8.179	4.312 -21.257	1.00	0.31
MOTA	444	CG LYS		8.954	5.631 -19.748	1.00	0.26
MOTA	445	HG1 LY		8.823			
MOTA	446	HG2 LY				1.00	0.40
				8.799	6.648 -19.430	1.00	0.42
ATOM	447	CD LY	33	10.380	5.469 -20.291	1.00	0.48
MOTA	448	HD1 LY	33	10.466	4.517 -20.793	1.00	0.74
MOTA	449	HD2 LY	33	11.080	5.505 -19.469	1.00	1.11
ATOM	450	CE LYS	3 33	10.705	6.593 -21.282	1.00	0.92
MOTA	451	HE1 LY		10.398	7.543 -20.868		
MOTA	452	HE2 LY	33	10.184		1.00	1.52
ATOM	453	NZ LY			6.419 -22.211	1.00	1.19
				12.172	6.614 -21.538	1.00	1.60
MOTA	454	HZ1 LY		12.668	6.957 -20.692	1.00	1.99
MOTA	455	HZ2 LY		12.374	7.247 -22.340	1.00	2.14
MOTA	456	HZ3 LY	33	12.498	5.653 -21.763	1.00	2.03
MOTA	457	C LY	3 3 3	6.399	4.158 -19.202	1.00	0.19
MOTA	458	O LY		6.054	3.035 -19.495		
ATOM	459	N AL				1.00	0.20
MOTA	460			6.682	4.471 -17.966	1.00	0.17
		HN AL		6.965	5.383 -17.740	1.00	0.18
MOTA	461	CA AL		6.589	3.428 -16.904	1.00	0.16
ATOM	462	HA AL	34	7.276	2.625 -17.128	1.00	0.18
ATOM	463	CB AL	34	6.952	4.043 -15.551	1.00	0.16
MOTA	464	HB1 AL		6.483	3.476 -14.761	1.00	1.02
ATOM	465	HB2 AL		6.604	5.065 -15.516		
ATOM	466	HB3 AL				1.00	0.98
MOTA	467		-	8.024	4.022 -15.423	1.00	1.02
		C AL		5.164	2.875 -16.844	1.00	0.16
ATOM	468	O AL		4.954	1.677 -16.847	1.00	0.17
MOTA	469	N PHI	35	4.182	3.729 -16.792	1.00	0.16
MOTA	470	HN PH	35	4.364	4.694 -16.792	1.00	0.16
ATOM	471	CA PHI		2.781	3.230 -16.736		
ATOM	472	HA PHI		2.690		1.00	0.17
MOTA	473	CB PH			2.525 -15.924	1.00	0.17
ATOM	474			1.815	4.396 -16.508	1.00	0.18
		HB1 PH		0.802	4.060 -16.672	1.00	0.19
MOTA	475	HB2 PH		2.045	5.192 -17.200	1.00	0.19
MOTA	476	CG PH		1.953	4.902 -15.089	1.00	0.18
MOTA	477	CD1 PH	35	1.616	4.071 -14.011	1.00	0.19
MOTA	478	HD1 PH	35	1.258	3.069 -14.191	1.00	0.19
MOTA	479	CD2 PH		2.415	6.203 -14.849	1.00	0.20
ATOM	480	HD2 PH	3 35	2.674	6.847 -15.677		
MOTA	481	CE1 PH	35	1.743	4 530 10 600	1.00	0.21
ATOM	482	HE1 PH			4.539 -12.699	1.00	0.21
MOTA	483			1.484	3.897 -11.870	1.00	0.23
				2.540	6.670 -13.535	1.00	0.22
MOTA	484	HE2 PH		2.893	7.672 -13.349	1.00	0.24
MOTA	485	CZ PH	3 35	2.205	5.838 -12.460	1.00	0.22
ATOM	486	HZ PH	35	2.303	6.198 -11.447	1.00	0.24
ATOM	487	C PHI		2.432	2.524 -18.048		
MOTA	488	O PH		1.770	1 507 -10 000	1.00	0.18
ATOM	489	N LY			1.507 -18.055	1.00	0.19
ATOM				2.864	3.053 -19.162	1.00	0.19
	490	HN LY		3.394	3.878 -19.144	1.00	0.19
MOTA	491	CA LY		2.535	2.399 -20.460	1.00	0.22
MOTA	492	HA LY		1.462	2.358 -20.574	1.00	0.23
MOTA	493	CB LY		3.135	3.205 -21.614	1.00	0.24
MOTA	494	HB1 LY		3.045	2.641 -22.530		
ATOM	495	HB2 LY	36	4.178		1.00	0.27
ATOM	496				3.400 -21.412	1.00	0.24
ATOM		CG LY		2.384	4.530 -21.758	1.00	0.27
	497	HG1 LY	36	2.471	5.097 -20.844	1.00	0.69
ATOM	498	HG2 LYS		1.341	4.332 -21.963	1.00	0.68
MOTA	499	CD LYS		2.988	5.332 -22.913	1.00	0.75
MOTA	50 <b>0</b>	HD1 LYS	36	2.898	4.766 -23.828	1.00	1.39
MOTA	501	HD2 LYS		4.032	5.525 -22.710		
				7.032	J.JEJ -42./1U	1.00	1.34

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3 0001	500								
MOTA	502		LYS	36	2.243	6.659	-23.065	1.00	1.15
MOTA	503	HE1	LYS	36	2.728	7.415	-22.464	1.00	
ATOM	504	HE2	LYS	36	1.221	6.540	-22.736		1.64
ATOM	505		LYS	36			-22./36	1.00	1.61
ATOM	506				2.260	7.076	-24.496	1.00	1.99
		HZ1		36	2.628	6.298	-25.079	1.00	2.51
MOTA	507		LYS	36	2.871	7.911	-24.605	1.00	2.40
ATOM	508	HZ3	LYS	36	1.295	7 300	-24.801		
ATOM	509		LYS	36	3.098	7.305	-24.601	1.00	2.38
ATOM	510						-20.481	1.00	0.21
			LYS	36	2.446	0.053	-20.927	1.00	0.23
MOTA	511	N	LYS	37	4.295	0.778	-19.995	1.00	0.21
ATOM	512	HN	LYS	37	4.810	1 527	-19.629	1.00	
ATOM	513		LYS	37	4.864	-0 600	10 000		0.20
ATOM	514		LYS			-0.600	-19.988	1.00	0.22
MOTA				37	4.926	-0.974	-21.000	1.00	0.24
	515		LYS	37	6.257	-0.581	-19.358	1.00	0.22
MOTA	516	HB1		37	6.589	-1.596	-19.195	1.00	0.24
ATOM	517	HB2	LYS	37	6.216	-0.061	-18.412		
ATOM	518		LYS	37	7.244	0.001	-10.412	1.00	0.21
MOTA	519	HG1				0.130	-20.285	1.00	0.26
		HGT.	1112	37	6.921	1.140	-20.459	1.00	0.25
MOTA	520	HG2		37	7.296	-0.398	-21.227	1.00	0.28
ATOM	521	CD	LYS	37	8.625	0.139	-19.628	1.00	
ATOM	522	HD1	LYS	37	8.994	-0.233	-19.551		0.30
ATOM	523	HD2		37		-0.673	-19.551	1.00	0.77
ATOM	524				8.549		-18.640	1.00	0.84
			LYS	37	9.594		-20.473	1.00	0.90
ATOM	525	HE1	LYS	37	10.530	1.076	-19.943	1.00	1.47
MOTA	526	HE2	LYS	37	9.169	1.945	-20.652	1.00	
ATOM	527		LYS	37	9.836	0.206	-20.052		1.59
ATOM	528	HZ1				0.286	-21.774	1.00	1.77
				37	9.798	0.984	-22.543	1.00	2.22
ATOM	529		LYS	37	9.106	-0.439	-21.926	1.00	2.28
MOTA	530	HZ3	LYS	37	10.774	-0.161	-21.762	1.00	
MOTA	531	C	LYS	37	3.955		10.150		2.33
MOTA	532		LYS	37			-19.158	1.00	0.20
ATOM					3.689	-2.636	-19.516	1.00	0.21
	533		ALA	38	3.479	-1.013	-18.046	1.00	0.19
MOTA	534	HN .	ALA	38	3.711	-0.098	-17 777	1.00	0.19
ATOM	535	CA .	ALA	38	2.589	-1.838	-17 102		
ATOM	536		ALA	38	3.116	2.000	-17.102	1.00	0.18
ATOM	537		ALA			-2.727	-16.870	1.00	0.19
				38	2.183	-1.030	-15.949	1.00	0.19
MOTA	538	HB1		38	2.831	-0.172	-15.851	1.00	1.05
MOTA	539	HB2	ALA	38	2.270	-1.649	-15.068	1.00	
ATOM	540	HB3	ALA	38	1.161	-0.698	-16 053		1.00
ATOM	541		ALA	38		-0.098	-10.027	1.00	1.06
MOTA	542				1.338	-2.238		1.00	0.18
			ALA	38	0.967	-3.392	-18.012	1.00	0.19
MOTA	543		PHE	39	0:688	-1.295	-18.589	1.00	0.18
ATOM	544	HN :	PHE	39 -	1.005	-0.368	-18 547		
MOTA	545	CA	PHE	39	-0.535	-1 633	-10.347	1.00	0.18
ATOM	546		PHE	39		-1.632	-19.367	1.00	0.19
MOTA	547				-1.248	-2.122	-18.720	1.00	0.19
			PHE	39 .	-1.156	-0.354	-19.937	1.00	0,21
MOTA	548		PHE	39	-1.883	-0.614	-20.692	1.00	0.24
ATOM	549	HB2	PHE	39	-0.381	0.256	-20.378		
ATOM	550	CG :	PHE	39	-1.836	0.230	-20.3/6	1.00	0.21
ATOM	551	CD1		39	-1.030	0.410	-18.829	1.00	0.20
ATOM	552				-3.010	-0.080	-18.250	1.00	0.25
		HD1		39	-3.429	-1.014	-18.595	1.00	0.30
ATOM	553	CD2	PHE	39	-1.294	1.627	-18.380	1.00	0.17
MOTA	554	HD2	PHE	39	-0.389	2 012	-18.827	1.00	
MOTA	555	CE1		39	-3.642	0.633	12.02/		0.18
ATOM	556		PHE	39		0.633	-17.224	1.00	0.28
MOTA	557				-4.548	0.250	-16.779	1.00	0.34
			PHE	39	-1.926	2.341	-17.354	1.00	0.18
ATOM	558		PHE	39	-1.507	3.275	-17.007	1.00	0.17
MOTA	559	CZ 1	PHE	39	-3.099		-16.776	1.00	0.27
MOTA	560	HZ 1	PHE	39	-3.587				0.23
ATOM	561	_	PHE	39		2.394	-15.985	1.00	0.26
ATOM	562	_			-0.154	-2.571	-20.508	1.00	0.18
			PHE	39	-0.862	-3.509	-20.817	1.00	0.18
ATOM	563		LYS	40	0.963	-2.330	-21 136	1.00	0.19
ATOM	564	HN 1	Lys	40	1.522	-1.570	-20 070		
ATOM	565		LYS	40	1 200	-4.5/0	~20.0/0	1.00	0.19
MOTA	566		LYS		1.388	-3.214	-22.254	1.00	0.19
MOTA				40	0.642	-3.186	-23.031	1.00	0.20
411.011	567		Lys	40	2.730	-2.707	-22.804	1.00	0.21
MOTA	568	HB1 I		40	3.466		-22.014	1.00	0.21
ATOM	569	HB2 I	LYS	40	2.610	-1.692	_22.014		
ATOM	570		LYS	40		-1.092	-23.155	1.00	0.25
ATOM	571				3.218	-3.588	-23.966	1.00	0.25
ATOM		HG1 I	223	40	3.337	-4.604 -	-23.621	1.00	0.46
	572	HG2 I		40	4.171	-3.218	-24.314	1.00	0.46
MOTA	573		LYS	40	2.213	-3.560	-25 121		
ATOM	574	HD1 I	LYS	40	1.840	_2.500 ·		1.00	0.38
ATOM	575	HD2 I	LYS	40		-2.555	-25.253	1.00	0.54
ATOM	576		YS	40	1.392	-4.227 -	-24.905	1.00	0.56
ATOM	577				2.903	-4.019 -	-26.407	1.00	0.40
		HE1 I		40	3.776	-4.604 -	-26.158	1.00	1.07
MOTA	578	HE2 I	712	40	3 199	-3 167 .	-24 OOE	1.00	1.07

ATOM	579	NZ	LYS	40	1.958	4 053	-27.203		
ATOM	580		LYS	-				1.00	1.40
			_	40	1.571	-5.607		1.00	1.95
ATOM	581	HZ2	LYS	40	2.464	-5.274	-28.009	1.00	1.92
ATOM	582	HZ3	LYS	40	1.181	-4.258	20.005		
								1.00	2.02
ATOM	583	C	LYS	40	1.553	-4.648	-21.740	1.00	0.17
ATOM	584	0	LYS	· 40	1.034	-5.583	-22.314	1.00	0.17
MOTA	585	N	VAL				-22.314		
				41	2.271	-4.828	-20.663	1.00	0.17
MOTA	586	HN	VAL	41	2.681	-4.060		1.00	0.18
MOTA	587	CA	VAL	41		6.000	20.214		
					2.468	-6.204	-20.116	1.00	0.16
MOTA	588	HA	VAL	41	2.953	-6.816	-20.862	1.00	0.17
ATOM	589	CB	VAL	41	3.350		10.002		
						-6.143		1.00	0.18
ATOM	590	HB	VAL	41	2.966	-5.393	-18.192	1.00	0.41
ATOM	591	CG1	VAL	41	3.343		-18.175		
ATOM	592					-7.308	-10.175	1.00	0.44
				41	2.420	-7.631	-17.629	1.00	1.16
MOTA	593	HG12	VAL	41	4.176	-7.571	-17.490		
ATOM	594					-7.571	-17.430	1.00	1.18
				41	3.429	-8.289	-18.916	1.00	1.11
ATOM	595		VAL	41	4.781	-5.785	-19.277	1.00	0.43
ATOM	596	HG21	MAT.	41					
				_	5.132	-6.492	-20.013	1.00	1.12
ATOM	597	HG22	VAL	41	5.423	-5.820	-18.411	1.00	1.11
MOTA	598	HG23	VAI.	41	4.797		-19.697		
ATOM						-4.790	-19.09/	1.00	1.19
	599	С	VAL	41	1.122	-6.833	-19.751	1.00	0.16
MOTA	600	0	VAL	41	0.887	-7 999	-19.996	1.00	0.17
ATOM	601	N	TRP	42					
					0.240	-6.080	-19.152	1.00	0.16
ATOM	602	HN	TRP	42	0.448	-5.143	-18.950	1.00	0.17
ATOM	603	CA	TRP	42		6 666	10.750		
					-1.079	-0.655	-18.761	1.00	0.17
ATOM	604	HA	TRP	42	-0.927	-7.642	-18.352	1.00	0.17
MOTA	605	CB	TRP	42	-1.739	5 767	17 600		
						~5./6/	-17.699	1.00	0.18
ATOM	606	HRI	TRP	42	-2.787	-6.018	-17.621	1.00	0.19
ATOM	607	HB2	TRP	42	-1.638	-4 730	-17.983		
ATOM	_					-4.730	-17.983	1.00	0.20
	608	CG	TRP	42	-1.073	-5.990	-16.377	1.00	0.18
MOTA	609	CD1	TRP	42	-0.311	-5 082	-15.724		
ATOM	610	HD1				-3.002	-13.724	1.00	0.22
				42	-0.092	-4.084	-16.066	1.00	0.28
ATOM	611	CD2	TRP	42	-1.095	-7 182	-15.539	1.00	0.19
ATOM	612	NE1	TRP			7.202	13.333		
				42	0.140	-5.643		1.00	0.22
ATOM	613	HE1	TRP	42	0.714	-5.194	-13.887	1.00	0.25
ATOM	614	CE2	TRP	42	-0.315	6 035	14 304		
						-0.935	-14.384	1:00	0.20
MOTA	615	CE3	TRP	42	-1.707	-8.441	-15.669	1.00	0.25
ATOM	616	HE3	TRP.	42	-2.309	0 650	-16.539		
ATOM						-8.038	-10.233	1.00	0.27
	617	CZ2	TRP	42	-0.149	-7.903	-13.393	1.00	0.24
ATOM	618	HZ2	TRP	42	0.454				
ATOM						-/.031	-12.521	1.00	0.25
	619	CZ3	TRP	42	-1.543	-9.418	-14.673	1.00	0.31
ATOM	620	HZ3	TRP	42	-2.018	-10 391	-14.782		
ATOM	621	CH2				-10.301	-14.702	1.00	0.39
			TRP	42	-0.764	-9.149	-13.538	1.00	0.30
ATOM	622	HH2	TRP	42	-0.642	-9.904	-12.775	1.00	0.35
ATOM	623	C.	TRP	42		3.304	-14.775		
					-1.991	-6.754	-19.985	1.00	0.17
MOTA	624	0	TRP	42	-2.726	-7.706	-20.138	1.00	0.18
ATOM	625	N	SER	43	-1.952	6 700	20.230		
ATOM						-5.782	-20.855	1.00	0.17
	626	HN	SER	43	-1.352	-5.021	-20.713	1.00	0.17
ATOM	627	CA	SER	43	-2.831	-E 02E	20 000		
ATOM	628					-3.623	-22.062	1.00	0.18
		HA	SER	43	-3.846	-6.028	-21.759	1.00	0.19
ATOM	629	CB	SER	43	-2.779	-4.474	-22.775		
ATOM	630	HB1						1.00	0.20
				43	-2.965		-22.059	1.00	0.21
ATOM	631	HB2	SER	43	-3.533	-4.442	-23.543	1.00	0.23
ATOM	632	OG	SER	43	-1.499	_4 304	22.273		
ATOM						-4.304	-23.368	1.00	0.21
	633	HG	SER	43	-1.031	-5.140	-23.309	1.00	0.97
ATOM	634	С	SER	43	-2.358	-6 022	-23.019		
MOTA	635	ō	SER			-0.322	-23.019	1.00	0.18
				43	-3.085	-/.350	-23.893	1.00	0.21
MOTA	636	N	ASP	44	-1.148		-22.866	1.00	0.17
ATOM	637	HN	ASP	44	-0.575				
ATOM							-22.156	1.00	0.18
	638	CA	ASP	44	-0.632	-8.445	-23.770	1.00	0.18
ATOM	639	HA	ASP	44	-0.650	-8 006	-24.788		
ATOM	640	CB				-0.000	-24./88	1.00	0.19
			ASP	44	0.809	-8.793	-23.386	1.00	0.20
MOTA	641	HB1	ASP	.44	1.117	-0 603	-23.915		
ATOM	642		ASP			2.003		1.00	0.21
				44	0.864	-8.969	-22.322	1.00	0.22
MOTA	643	CG	ASP	44	1.734		-23.760		
MOTA	644		ASP					1.00	0.24
				44	1.340	-6.833	-24.591	1.00	0.85
MOTA	645	OD2	ASP	44	2.820	-7.568	-23.209	1.00	0.84
MOTA	646	C	ASP	44					
					-1.499		-23.665	1.00	0.19
MOTA	647	0	ASP	44	-1.753	-10.366		1.00	0.21
MOTA	648	N	VAL	45	-1.927				
ATOM								1.00	0.21
	649	HN	VAL	45	-1.689	-9.519	-21.693	1.00	0.21
ATOM	650	CA	VAL	45		-11.299	22 202		
ATOM	651	HA	VAL		-6./93	-11.639	-22.302	1.00	0.26
				45	-2.833	-11.811	-23.247	1.00	0.28
ATOM	652	CB	VAL	45		-12.222	-21 303		
MOTA	653	HB	VAL	45	2.043	-14.666	-21.303	1.00	0.30
					-2.645	-13.107	-21.146	1.00	0.37
MOTA	654		VAL	45	-0.678	-12.626	-21.866	1.00	0.36
MOTA	655	HG11	VAI.	45	_0.010	11 77	22.000		
				-3	-0.210	-11 766	-33 333	1 00	4 //7

	474 11546 411-			
MOTA	656 HG12 VAL	45	-0.810 -13.400 -22.607 1.00 1.02	
MOTA	657 HG13 VAL	45	-0.051 -12.995 -21.068 1.00 1.13	
MOTA	658 CG2 VAL	45	-1.855 -11.486 -19.973 1.00 0.32	
MOTA	659 HG21 VAL	45	-2.819 -11.303 -19.524 1.00 0.96	
ATOM	660 HG22 VAL	45		
			-1.356 -10.545 -20.149 1.00 1.09	
MOTA	661 HG23 VAL	45	-1.258 -12.091 -19.305 1.00 1.11	
ATOM	662 C VAL	45	-4.160 -10.966 -21.790 1.00 0.29	
MOTA	663 O VAL	45	-4.837 -11.819 -21.249 1.00 0.64	
MOTA	664 N THR	46	-4.619 -9.748 -21.963 1.00 0.36	
ATOM	665 HN THR	46		
			-4.062 -9.076 -22.409 1.00 0.65	
MOTA	666 CA THR	46	-5.998 -9.382 -21.491 1.00 0.38	
MOTA	667 HA THR	46	-6.567 -10.277 -21.320 1.00 0.44	
ATOM	668 CB THR	46	-5.912 -8.577 -20.186 1.00 0.39	
MOTA	669 HB THR	46		
ATOM	670 OG1 THR			
	:	46	-5.018 -7.491 -20.358 1.00 0.36	
ATOM	671 HG1 THR	46	-5.532 -6.719 -20.608 1.00 0.94	
MOTA	672 CG2 THR	46	-5.430 -9.461 -19.036 1.00 0.43	
MOTA	673 HG21 THR	46	-4.929 -10.327 -19.429 1.00 1.08	
MOTA	674 HG22 THR	46	-6.277 -9.775 -18.445 1.00 1.15	
MOTA	675 HG23 THR	46		
ATOM	676 C THR	46	-6.668 -8.482 -22.553 1.00 0.32	•
MOTA	677 O THR	46	-6.124 -7.450 -22.892 1.00 0.32	
MOTA	678 N PRO	47	-7.833 -8.829 -23.084 1.00 0.30	
ATOM	679 CA PRO	47	-8.479 -7.951 -24.100 1.00 0.30	
ATOM	680 HA PRO	47		
MOTA				
		47	-9.687 -8.773 -24.546 1.00 0.35	
ATOM	682 HB1 PRO	47	-9.541 -9.110 -25.561 1.00 0.40	
MOTA	683 HB2 PRO	47	-10.579 -8.166 -24.489 1.00 0.37	
MOTA	684 CG PRO	47	-9.825 -9.986 -23.621 1.00 0.35	
ATOM	685 HG1 PRO	47	-9.916 -10.885 -24.212 1.00 0.42	
ATOM	686 HG2 PRO	47		
MOTA	687 CD PRO	47	-8.576 -10.077 -22.739 1.00 0.33	
MOTA	688 HD2 PRO	47	-8.853 -10.091 -21.692 1.00 0.31	
MOTA	689 HD1 PRO	47	-7.993 -10.946 -22.999 1.00 0.39	
ATOM	690 C PRO	47	-8.933 -6.614 -23.506 1.00 0.25	
ATOM	691 O PRO	47	-9.744 -5.914 -24.080 1.00 0.26	
MOTA	692 N LEU	48		
ATOM				
		48	-7.766 -6.828 -21.912 1.00 0.29	
MOTA	694 CA LEU	48	-8.827 -4.960 -21.742 1.00 0.26	
MOTA	695 HA LEU	48	-9.904 -4.905 -21.696 1.00 0.27	
ATOM	696 CB LEU	48	-8.241 -4.858 -20.329 1.00 0.31	
MOTA	697 HB1 LEU	48	-8.476 -3.892 -19.909 1.00 0.34	
ATOM	698 HB2 LEU	48	-7.167 -4.968 -20.385 1.00 0.33	
ATOM	699 CG LEU	48		
ATOM			-8.816 -5.964 -19.434 1.00 0.34	
		48	-8.808 -6.900 -19.972 1.00 0.32	
MOTA	701 CD1 LEU	48	-7.952 -6.091 -18.177 1.00 0.41	
ATOM	702 HD11 LEU	48	-8.002 -5.171 -17.613 1.00 1.11	
ATOM	703 HD12 LEU	48	-6.928 -6.283 -18.462 1.00 1.05	
MOTA	704 HD13 LEU	48	-8.315 -6.906 -17.570 1.00 1.15	
ATOM	705 CD2 LEU	48		
ATOM	706 HD21 LEU	-	71.71	
		48	-10.569 -4.707 -19.478 1.00 1.10	
ATOM	707 HD22 LEU	48	-10.299 -5.524 -17.942 1.00 1.09	
MOTA	708 HD23 LEU	48	-10.912 -6.428 -19.325 1.00 1.04	
MOTA	709 C LEU	48	-8.289 -3.806 -22.589 1.00 0.25	
ATOM	710 O LEU	48	-7.174 -3.849 -23.071 1.00 0.26	
MOTA	711 N ASN	49	-9.073 -2.775 -22.762 1.00 0.25	
MOTA	712 HN ASN	49		
ATOM			-9.964 -2.770 -22.355 1.00 0.26	
		49	-8.622 -1.604 -23.568 1.00 0.25	
MOTA	714 ha asn	49	-7.703 -1.842 -24.082 1.00 0.27	
ATOM	715 CB ASN	49	-9.700 -1.245 -24.593 1.00 0.28	
MOTA	716 HB1 ASN	49	-9.390 -0.375 -25.153 1.00 0.30	
ATOM	717 HB2 ASN	49	-10.628 -1.033 -24.081 1.00 0.28	
MOTA	718 CG ASN	49		
ATOM				
		49	-9.798 -3.564 -25.161 1.00 1.10	
MOTA	720 ND2 ASN	49	-10.186 -2.182 -26.804 1.00 1.14	
MOTA	721 HD21 ASN	49	-10.268 -1.258 -27.121 1.00 1.94	
MOTA	722 HD22 ASN	49	-10.317 -2.927 -27.427 1.00 1.14	
MOTA	723 C ASN	49	-8.391 -0.417 -22.633 1.00 0.24	
MOTA	724 O ASN	49	-9.290 0.016 -21.939 1.00 0.23	,
ATOM	725 N PHE	50		
			-7.192 0.107 -22.606 1.00 0.24	
ATOM	726 HN PHE	50	-6.485 -0.264 -23.173 1.00 0.26	
MOTA	727 CA PHE	50	-6.896 1.263 -21.710 1.00 0.23	j
MOTA	728 HA PHE	50	-7.688 1.380 -20.985 1.00 0.21	
ATOM	729 CB PHE	50	-5.574 1.016 -20.981 1.00 0.24	
ATOM	730 HB1 PHE	50	-5.357 1.853 -20.334 1.00 0.25	
MOTA	731 HB2 PHE	50		,
ATOM	732 CG PHE	50		
	co inc	.,,,	-5 676 -0 743 -20 154 1 nn n 23	

ATOM	733	CD1	PHE	50	-6.266	-0.201 -18.886	1 00	0.25
ATOM	734	HD1	PHE	50	-6.652	0.731 -18.500	1.00	
ATOM	735	CD2	PHE	50			1.00	0.28
ATOM	736				-5.176	-1.451 -20.654	1.00	0.22
			PHE	50	-4.720	-1.483 -21.633	1.00	0.23
ATOM	737		PHE	50	-6.358	-1.368 -18.117	1.00	0.25
MOTA	738	HE1	PHE	50	-6.813	-1.336 -17.139	1.00	0.28
ATOM	739	CE2	PHE	50	-5.267	-2.618 -19.886	1.00	0.23
ATOM	740	HE2	PHE	50	-4.881	-3.550 -20.272		
ATOM	741	CZ					1.00	0.25
			PHE	50	-5.858	-2.576 -18.618	1.00	0.24
ATOM	742	HZ	PHE	50	-5.928	-3.476 -18.025	1.00	0.25
ATOM	743	C	PHE	50	-6.777	2.538 -22.545	1.00	0.26
MOTA	744	0	PHE	50	-6.028	2.596 -23.501	1.00	0.31
MOTA	745	N	THR	51	-7.517	3.555 -22.184	1.00	0.24
ATOM.	746	HN	THR	51	-8.109	3.468 -21.413		
ATOM	747	CA	THR	51	-0.109	3.468 -21.413	1.00	0.22
					-7.470	4.842 -22.940	1.00	0.27
ATOM	748	HA	THR	51	-6.775	4.762 -23.762	1.00	0.31
MOTA	749	CB	THR	51	-8.868	5.153 -23.483	1.00	0.30
MOTA	750	HB	THR	51	-9.562	5.248 -22.663	1.00	0.29
MOTA	751	OG1	THR	51	-9.283	4.100 -24.341	1.00	0.35
MOTA	752	HG1	THR	51	-9.638	4.491 -25.142		
MOTA	753	CG2	THR	51		6 464 24 252	1.00	0.84
ATOM		HG21			-8.835	6.464 -24.273	1.00	0.34
			THR	51	-9.805	6.640 -24.716	1.00	1.02
ATOM	755	HG22	THR	51	-8.092	6.394 -25.053	1.00	1.07
ATOM	756	HG23	THR	51	-8.588	7.280 -23.611	1.00	1.13
MOTA	757	C	THR	51	-7.024	5.969 -22.001	1.00	0.25
ATOM	758	0	THR	51	-7.553	6.139 -20.920		
MOTA	759	N	ARG	52	-6.054	6.740 20.920	1.00	0.22
ATOM	760					6.740 -22.411	1.00	0.29
		HN	ARG	52	-5.645	6.583 -23.287	1.00	0.32
ATOM	761	CA	ARG	52	-5.566	7.861 -21.556	1.00	0.29
ATOM	762	HA	ARG	52	-5.591	7.563 -20.518	1.00	0.27
MOTA	763	CB	ARG	52	-4.128	8.201 -21.955	1.00	0.35
ATOM	764	HB1	ARG	52	-4.125	8.654 -22.935		
ATOM	765		ARG	52	-3.539	7 205 21 25	1.00	0.39
ATOM	766	CG				7.295 -21.977	1.00	0.38
		-	ARG	52	-3.521	9.177 -20.945	1.00	0.39
MOTA	767		ARG	52	-3.645	8.787 -19.946	1.00	0.71
MOTA	768		ARG	52	-4.017	10.134 -21.025	1.00	0.57
MOTA	769	CD	ARG	52	-2.030	9.345 -21.244	1.00	0.79
MOTA	770	HD1	ARG	52	-1.825	9.001 -22.248		
ATOM	771		ARG	52		9.001 -22.240	1.00	1.45
ATOM	772	NE			-1.453	8.763 -20.543	1.00	1.39
			ARG	52	-1.656	10.782 -21.120	1.00	1.47
ATOM	773	HE	ARG	52	-2.354	11.468 -21.073	1.00	2.06
MOTA	774	CZ	ARG	52	-0.398	11.127 -21.071	1.00	2.09
ATOM	775	NH1	ARG	52	-0.070	12.385 -20.960	1.00	3.05
MOTA	776	HH11	ARG	52	-0.782	13.084 -20.911		3.05
ATOM	777	HH12		52	0.894		1.00	3.45
ATOM	778		ARG			12.649 -20.923	1.00	3.60
ATOM				52	0.532	10.213 -21.138	1.00	2.31
		HH21		52	0.281	9.249 -21.226	1.00	2.16
MOTA		HH22		52	1.496	10.477 -21.102	1.00	3.05
ATOM	781	C	ARG	52	-6.460	9.090 -21.758	1.00	0.29
MOTA	782	0	ARG	52	-6.719	9.495 -22.875	1.00	0.33
ATOM	783	N	LEU	53	-6.928	9.689 -20.689		
MOTA	784	HN		53	-0.320	9.689 -20.689	1.00	0.26
ATOM	785				-6.702	9.345 -19.798	1.00	0.25
		CA	LEU	53	-7.803	10.896 -20.822	1.00	0.29
MOTA	786	HA	LEU	53	-8.167	10.972 -21.835	1.00	0.32
MOTA	787	CB	LEU	53	-8.992	10.784 -19.862	1.00	0.28
MOTA	788	HB1	LEU	53	-9.579	11.688 -19.908	1.00	0.31
MOTA	789	HB2	LEU	53	-8.624	10.648 -18.855	1.00	0.28
MOTA	790	CG	LEU	53	-9.866	9.587 -20.249		
MOTA	791	HG	LEU	53		9.507 -20.245	1.00	0.28
ATOM	792		LEU		-9.264	8.690 -20.246	1.00	0.29
ATOM		1201	LEU	53	-10.999	9.440 -19.232	1.00	0.29
	793	HD11	TEO	53	-11.606	8.585 -19.487	1.00	0.95
MOTA	794	HD12	LEU	53	-11.610	10.331 -19.243	1.00	1.05
MOTA	795	HD13	LEU	53	-10.581	9.303 -18.247	1.00	1.07
ATOM	796	CD2	LEU	53	-10.463	9.799 -21.646		
ATOM		HD21	LEII	53		10 055 05 050	1.00	
ATOM	700	HD22	LETT		-10.523	10.856 -21.860	1.00	1.01
	700	UD22		53	-11.453	9.370 -21.685	1.00	1.09
MOTA		HD23		53	-9.835	9.319 -22.382	1.00	1.14
ATOM	800	C	LEU	53	-7.000	12.154 -20.483	1.00	0.33
MOTA	801	0	LEU	53	-6.315	12.218 -19.482	1.00	0.34
MOTA	802		HIS	54	-7.080	13.154 -21.319		
ATOM	803	HN	HIS	54		12 075 00 100	1.00	0.41
ATOM	804	CA	HIS		-7.637	13.075 -22.121	1.00	0.45
ATOM				54	-6.324	14.413 -21.062	1.00	0.47
	805	HA	HIS	54	-5.292	14.183 -20.851	1.00	0.54
MOTA	806	CB	HIS	54	-6.407	15.314 -22.297	1.00	0.60
MOTA	807	HB1	HIS	54	-6.018	16.291 -22.052	1.00	0.64
MOTA	808	HB2	HIS	54	-7.438	15.407 -22.603	1.00	0.61
ATOM	809	CG	HIS	54	-5.602			
-					-3.002	14.726 -23.426	1.00	0.74

ATOM	810	ND1	HTC	54	-5.645	15 254	-24.707		
								1.00	1.35
MOTA	811	HD1		54	-6.172		-24.996	1.00	1.86
MOTA	812	CD2	HIS	54	-4.740	13.656	-23.493	1.00	0.86
MOTA	813	HD2	HTS	54	-4.480		-22.668		
ATOM								1.00	1.34
	814	CE1		54	-4.834	14.512	-25.481	1.00	1.33
ATOM	815	HE1	HIS	54	-4.670	14.692	-26.533	1.00	1.83
ATOM	816	NE2		54	-4.257		-24.792		
								1.00	0.92
MOTA	817	C	HIS	54	-6.933	15.154	-19.867	1.00	0.43
ATOM	818	0	HIS	54	-6.230	15.714	-19.051	1.00	0.49
MOTA	819		ASP	55					
					-8.236	15.172	-19.767	1.00	0.42
ATOM	820	HN	ASP	55	-8.784	14.719	-20.442	1.00	0.45
ATOM.	821	CA	ASP	55	-8.892	15.892	-18.635	1.00	0.49
MOTA	822			55			-10.033		
			ASP		-8.217	15.938	-17.796	1.00	0.54
MOTA	823	CB	ASP	<b>.</b> 55	-9.251	17.314	-19.073	1.00	0.65
MOTA	824	HB1	ASP	` 55	-9.876		-18.323		
MOTA	825						-10.323	1.00	0.75
		HB2		55	-9.783	17.277	-20.013	1.00	0.68
MOTA	826	CG	ASP	55	-7.974	18.140	-19.244	1.00	0.71
MOTA	827	OD1	ASP	55	-7.978	19.037	-20.071		
MOTA	828							1.00	1.19
		OD2		55	-7.018		-18.536	1.00	1.28
ATOM	829	C	ASP	55	-10.167	15.156	-18.223	1.00	0.45
MOTA	830	0	ASP	55	-10.638		-18.912		
ATOM	831					14.4/3	-10.912	1.00	0.44
			GLY	56	-10.728		-17.100	1.00	0.46
MOTA	832	HN	GLY	56	-10.328	16.233	-16.563	1.00	0.50
ATOM	833	CA	GLY	56	-11.975	14.848	-16.632	1.00	
MOTA	834	HA1		56			-10.032		0.44
					-12.482	14.399	-17.472	1.00	0.44
ATOM	835		GLY	56	-12.622	15.579	-16.169	1.00	0.48
ATOM	836	С	GLY	56	-11.624	13.760	-15.614		
ATOM	837					13.700	-13.014	1.00	0.40
			GLY	56	-10.473	13.543	-15.294	1.00	0.42
MOTA	838	N	ILE	57	-12.613	13.078	-15.105	1.00	0.37
ATOM	839	HN	ILE	57	-13.533				
ATOM					-13.333		-15.380	1.00	0.39
_	840	CA	ILE	57	-12.352	12.002	-14.106	1.00	0.35
ATOM	841	HA	ILE	57	-11.406		-13.616	1.00	0.38
ATOM	842	CB	ILE	57	-13.473				
ATOM							-13.064	1.00	0.41
	843	HB	ILE	57	-14.415	11.820	-13.561	1.00	0.42
ATOM	844	CG1	ILE	57	-13.508	13.363	-12.360	1.00	0.48
MOTA	845	HG11	ILE	57	-13.512				
						:	-13.101	1.00	0.48
ATOM	846		ILE	57	-12.631	13.465	-11.737	1.00	0.51
ATOM	847	CG2	ILE	57	-13.216	10.896	-12.037	1.00	0.44
ATOM	848		ILE	57	-13.315	0.030	12.037		
	-					9.932	-12.513	1.00	1.19
MOTA	849		ILE	57	-13.934	10.977	-11.235	1.00	1.09
ATOM	850	HG23	ILE	57	-12.218		-11.639	1.00	1.04
MOTA	851		ILE	57					
					-14.765		-11.488	1.00	0.56
MOTA	852	HD11	ILE	57	-15.459	12.693	-11.728	1.00	1.08
ATOM	853	HD12	ILE	57	-15.235	14.439	-11.668		
ATOM	854	HD13	ILE	57				1.00	1.24
				••	-14.487	13.413	-10.447	1.00	1.14
ATOM	855	C	ILE	57	-12.307	10.647	-14.817	1.00	0.30
ATOM	856	0	ILE	57	-13.139		-15.653	1.00	0.31
MOTA	857		ALA	58	-11.337		-13.033		
						9.828	-14.493	1.00	0.26
ATOM	858	HN	ALA	58	-10.679	10.096	-13.817	1.00	0.27
ATOM	859	CA	ALA	58	-11.221	8.489	-15.148	1.00	0.23
ATOM	860	HA	ALA	58	-11.957		15.110		
ATOM					-11.33/	8.398	-15.932	1.00	0.25
	861		ALA	58	-9.824	8.339	-15.749	1.00	0.23
ATOM	862		ALA	58	-9.843	7.585	-16.522	1.00	0.97
MOTA	863	HB2	ALA	58	-9.129		-14.976	1 00	
ATOM	864		ALA			0.024	-13.7/0	1.00	1.11
				58	-9.513	9.280	-16.172	1.00	1.03
ATOM	865	C	ALA	58	-11.443	7.387	-14.114	1.00	0.23
ATOM	866	0	ALA	58	-11.389		-12.922	1.00	0.27
MOTA	867		ASP	59	-11.701				
ATOM							-14.564	1.00	0.25
	868		ASP	59	-11.744	6.028	-15.530	1.00	0.28
MOTA	869	CA	ASP	59	-11.934	5.069	-13.613	1.00	0.27
ATOM	870		ASP	59	-12.788	E 2003	12.013		
ATOM						2.496	-12.991	1.00	0.34
	871		ASP	59	-12.207	3.785	-14.400	1.00	0.33
MOTA	872	HB1	ASP	59	-12.203	2.942	-13.725	1.00	0.34
MOTA	873		ASP	59		2 276	-10.743		
MOTA					-11.438	2.051	-15.147	1.00	0.32
	874		ASP	59	-13.572	3.880	-15.084	1.00	0.44
MOTA	875	OD1	ASP	59	-13.791	3 130	-16.028	1.00	
MOTA	876	OD2				4.139	-10.028		1.20
				59	-14.374	4.691	-14.653	1.00	1.14
ATOM	877		ASP	59	~10.700	4.863	-12.731	1.00	0.22
MOTA	878	0	ASP	59	-10.806	A 767	-11.524		
MOTA	879	N	ILE			4.70/	-11.324	1.00	0.27
				60	-9.534	4.780	-13.326	1.00	0.18
MOTA	880	HN	ILE	60	-9.478	4.850	-14.302	1.00	0.20
ATOM	881	CA	ILE	60	-8.291	4 551	-12 502	1 00	
ATOM	882	HA	ILE			4.301	-12.523	1.00	0.22
				60	-8.554	4.303	-11.512	1.00	0.28
MOTA	883	CB	ILE	60	-7.502	3.404	-13.155	1.00	0.27
MOTA	884	HB	ILE	60	-7.255	3 655	-14.175		
ATOM	885	CG1				2.033	-14.1/3	1.00	0.28
	000	COT	TUE	60	-8.377	2.146	-13.136	1.00	N. 30

MOTA		HG12		60	-8.541	1.839	-12.113	1.00	0.36
MOTA	888	CG2	ILE	60	-6.210	3.127	-12.369	1.00	0.39
MOTA	889	HG21	ILE	60	-6.456	2.704	-11.409	1.00	1.05
MOTA	890	HG22	ILE	60	-5.658		-12.228	1.00	1.10
ATOM	891	HG23	ILE	60	-5.600		-12.921	1.00	1.12
ATOM	892	CD1		60	-7.688		-13.904	1.00	0.38
ATOM			ILE	60	-7.209		-14.786	1.00	1.07
ATOM			ILE	60	-8.424				
		HD13	ILE	60		0.280	-14.196	1.00	1.14
MOTA					-6.948		-13.270	1.00	1.04
MOTA	896	C	ILE	60	-7.438		-12.518	1.00	0.20
ATOM	897	0	ILE	60	-6.731	6.115	-13.464	1.00	0.25
MOTA	898	N	MET	61	-7.473		-11.448	1.00	0.20
MOTA	899	HN	MET	61	~8.033		-10.687	1.00	0.25
MOTA	900	CA	MET	61	-6.641		-11.373	1.00	0.20
MOTA	901	HA	MET	61	-6.327	8.102	-12.366	1.00	0.19
MOTA	902	CB	MET	61	-7.464		-10.773	1.00	0.24
ATOM	903	HB1	MET	61	-8.331	9.137	-11.392	1.00	0.35
MOTA	904	HB2	MET	61	-6.860	9.856	-10.743	1.00	0.33
ATOM	905	CG	MET	61	-7.918	8.604	-9.358	1.00	0.31
ATOM	906	HG1		61	-7.146	8.870	-8.653	1.00	0.66
ATOM	907	HG2		61	-8.112	7.544	-9.300	1.00	0.67
ATOM	908	SD	MET	61	-9.433	9.519	-8.967	1.00	0.54
ATOM	909	CE	MET	61	-8.878	11.154	-9.516		0.40
ATOM	910	HE1		61	-9.492			1.00	
ATOM	911	HE2	MET		-8.968	11.914	-9.056	1.00	1.06
	912			61			-10.589	1.00	1.16
MOTA		HE3	MET	61	-7.846	11.298	-9.232	1.00	1.12
MOTA	913	Ç	MET	61	-5.396		-10.524	1.00	0.20
ATOM	914	0	MET	61	-5.478	6.951	-9.463	1.00	0.22
MOTA	915	N	ILE	62	-4.241		-11.001	1.00	0.20
MOTA	916	HN	ILE	62	-4.207		-11.868	1.00	0.21
ATOM	917	CA	ILE	62	-2.971	7.678	-10.252	1.00	0.21
MOTA	918	HA	ILE	62	-3.156	6.982	-9.448		0.20
MOTA	919	CB	ILE	62	-1.938	7.080	-11.211	1.00	0.24
MOTA	920	HB	ILE	62	-1.753	7.781	-12.012	1.00	0.26
MOTA	921	CG1		62	-2.480	5.762	-11.785	1.00	0.23
ATOM	922	HG11	ILE	62	-3.479		-12.162		0.20
MOTA	923		ILE	62	-2.508		-11.003	1.00	
ATOM	924	CG2		62				1.00	0.24
ATOM		HG21		62	-0.635		-10.455	1.00	0.30
					-0.863	6.443	-9.466	1.00	1.08
ATOM		HG22	ILE	62	-0.070	7.729	-10.375	1.00	1.12
ATOM	927		ILE	62	-0.052	6.076	-10.988	1.00	0.99
ATOM	928	CD1		62	-1.584		-12.927	1.00	0.29
ATOM		HD11	ILE	62	-0.979		-13.305	1.00	1.02
ATOM		HD12	ILE	62	-2.201		-13.724	1.00	1.09
MOTA	931	HD13	ILE	62	-0.941	4.476	-12.559	1.00	1.07
ATOM	932	C	ILE	62	-2.423	8.988	-9.677	1.00	0.22
MOTA	933	0	ILE	62	-2.393	10.004	-10.343	1.00	0.27
MOTA	934	N	SER	63	-1.993	8.976	-8.441	1.00	0.20
MOTA	935	HN	SER	63	-2.028	8.147	-7.916	1.00	0.18
MOTA	936	CA	SER	63	-1.452	10.226	-7.829	1.00	0.22
MOTA	937	HA	SER	63	-0.998	10.836	-8.597	1.00	0.26
MOTA	938	CB	SER	63	-2.597	11.000	-7.176	1.00	0.24
ATOM	939		SER	63	-3.448	11.012	-7.845	1.00	0.25
MOTA	940		SER	63	-2.286	12.012	-6.978	1.00	0.29
MOTA	941	OG	SER	63	-2.951	10.369	-5.952	1.00	0.25
MOTA	942	HG	SER	63	-3.682	9.772			
ATOM	943	Ċ	SER	63	-0.404		-6.127 -6.764	1.00	0.85
ATOM	944	ŏ	SER	63		9.879		1.00	0.21
MOTA	945	Ŋ	PHE		-0.364	8.775	-6.259	1.00	0.20
				64	0.440	10.823	-6.419	1.00	0.24
ATOM	946	HN	PHE	64	0.380	11.705	-6.841	1.00	0.27
MOTA	947	CA	PHE	64 .	1.490	10.569	-5.382	1.00	0.24
ATOM	948	HA	PHE	64	1.560	9.511	-5.179	1.00	0.22
MOTA	949	CB	PHE	64	2.840	11.084	-5.895	1.00	0.28
MOTA	950	HB1		64	3.564	11.047	-5.097	1.00	0.32
MOTA	951	HB2	PHE	64	2.730	12.103	-6.235	1.00	0.32
MOTA	952	CG	PHE	64	3.316	10.220	-7.040	1.00	0.28
MOTA	953	CD1	PHE	- 64	4.112	9.096	-6.788	1.00	0.30
MOTA	954		PHE	64	4.385	8.844	-5.774	1.00	0.32
ATOM	955		PHE	64	2.963	10.545	-8.355	1.00	0.33
ATOM	956			64	2.350	11.412	-8.550	1.00	0.37
ATOM	957		PHE	64	4.553	8.297			
ATOM	958		PHE	64	5.166		-7.850	1.00	0.36
MOTA	959	CE2	PHE	64		7.430	-7.656	1.00	0.40
ATOM	960	HE2	PHE		3.403	9.747	-9.417	1.00	0.40
ATOM	961	CZ		64	3.130		-10.431	1.00	0.47
			PHE	64	4.198	8.623	-9.165	1.00	0.40
MOTA	962	HZ	PHE	64	4.538	8.007	-9.984	1.00	0.47
MOTA	963	С	PHE	64	1.115	11.318	-4.097	1.00	0.27

MOTA	964	0	PHE	64	0.924	12.518	-4.108	1.00	0.36
ATOM	965	N	GLY	65	0.996	10.617	-2.996	1.00	0.30
ATOM	966	HN	GLY	65	1.146	9.649	-3.017		
	967			65	– – –			1.00	0.33
ATOM		CA	GLY		0.615	11.282	-1.709	1.00	0.38
ATOM	968		GLY	65	-0.152	10.697	-1.224	1.00	0.46
ATOM	969	HA2	GLY	65	0.230	12.270	-1.913	1.00	0.45
MOTA	970	С	GLY	65	1.823	11.397	-0.770	1.00	0.32
MOTA	971	0	GLY	65	2.926	11.007	-1.098	1.00	0.40
ATOM	972	N	ILE	66	1.598	11.926			
							0.408	1.00	0.30
MOTA	973	HN	ILE	66	0.691	12.220	0.635	1.00	0.36
MOTA	974	CA	ILE	66	2.691	12.081	1.417	1.00	0.36
MOTA	975	HA	ILE	66	3.564	11.534	1.093	1.00	0.40
MOTA	976	CB	ILE	66	3.040	13.564	1.571	1.00	0.41
ATOM	977	HB	ILE	66	2.127	14.134	1.656	1.00	0.64
ATOM	978	CG1		66	3.829	14.026			
		HG11					0.337	1.00	0.68
ATOM			ILE	66	3.301	13.729	-0.557	1.00	0.95
MOTA	980	HG12	ILE	66	4.804	13.561	0.346	1.00	1.01
ATOM	981	CG2	ILE	66	3.886	13.764	2.831	1.00	0.93
MOTA	982	HG21	ILE	66	4.372	14.727	2.790	1.00	1.50
ATOM	983	HG22	ILE	66	4.632	12.986	2.891	1.00	1.41
ATOM	984	HG23	ILE	66	3.249	13.720	3.702		
	985							1.00	1.54
ATOM		CD1	ILE	66	3.997	15.551	0.343	1.00	0.70
ATOM		HD11	ILE	66	4.944	15.806	0.797	1.00	1.22
ATOM	987	HD12	ILE	66	3.196	16.009	0.902	1.00	1.28
ATOM	988	HD13	ILE	66	3.979	15.917	-0.673	1.00	1.23
ATOM	989	С	ILE	66	2.207	11.519	2.760	1.00	0.46
MOTA	990	ŏ	ILE	66	1.021	11.363	2.750		
MOTA	991							1.00	0.54
		N	LYS	67	3.129	11.205	3.659	1.00	0.59
MOTA	992	HN	LYS	67	4.073	11.343	3.434	1.00	0.64
MOTA	993	CA	LYS	67	2.780	10.630	5.014	1.00	0.74
MOTA	994	HA	LYS	67	3.072	9.594	5.038	1.00	0.83
ATOM	995	CB	LYS	67	3.550	11.404	6.102	1.00	
ATOM	996		LYS	67	3.237				0.90
						12.438	6.089	1.00	0.89
MOTA	997		LYS	67	4.608	11.352	5.891	1.00	0.96
MOTA	998	CG	LYS	67	3.287	10.815	7.504	1.00	1.08
MOTA	999	HG1	LYS	67	2.254	10.524	7.598	1.00	1.31
MOTA	1000	HG2	LYS	67	3.510	11.565	8.249	1.00	1.33
ATOM	1001	CD	LYS	· 67	4.179	9.590	7.746	1.00	0.98
ATOM	1002		LYS	67	5.216				
ATOM	1003					9.885	7.694	1.00	1.07
			LYS	67	3.979	8.839	6.999	1.00	1.07
ATOM	1004	CE	LYS	67	3.885	9.016	9.135	1.00	1.17
ATOM	1005	HE1	LYS	67	4.331	8.036	9.220	1.00	1.64
ATOM	1006	HE2	LYS	67	2.817	8.938	9.272	1.00	1.50
ATOM	1007	NZ	LYS	67	4.453	9.913	10.180	1.00	1.93
ATOM	1008		LYS	67	4.569	10.870			
ATOM	1009	HZ2	LYS				9.792	1.00	2.38
				67	5.378	9.547	10.485	1.00	2.43
ATOM	1010	HZ3	LYS	67	3.808	9.948	10.995	1.00	2.40
MOTA	1011	C	LYS	67	1.274	10.732	5.280	1.00	0.72
MOTA	1012	0	LYS	67	0.530	9.804	5.035	1.00	0.79
MOTA	1013	N	GLU	68	0.815	11.855	5.760	1.00	0.77
ATOM	1014	HN	GLU	68	1.425				
MOTA	1015	CA	GLU			12.601	5.939	1.00	0.84
				68	-0.645	12.004	6.011	1.00	0.84
MOTA	1016	HA	GLU	68	-1.014	11.130	6.530	1.00	0.99
MOTA	1017	CB	GLU	68	-0.895	13.254	6.860	1.00	1.05
MOTA	1018	HB1		68	-0.393	13.149	7.810	1.00	1.23
MOTA	1019	HB2	GLU	68	-1.956	13.370	7.024	1.00	1.10
ATOM	1020	CG	GLU	68	-0.353	14.487	6.134	1.00	1.15
MOTA	1021		GLU	68	-1.000	14.730	5.304	1.00	1.32
ATOM	1022	HG2	GLU	68	0.642				1.32
ATOM	1023					14.281	5.768	1.00	1.28
		CD	GLU	68	-0.308	15.669	7.104	1.00	1.75
MOTA	1024	OE1	GLU	68	0.246	16.692	6.736	1.00	2.45
MOTA	1025	OE2	GLU	68	-0.823	15.530	8.202	1.00	2.16
MOTA	1026	С	GLU	68	-1.346	12.132	4.660	1.00	0.76
MOTA	1027	Ō	GLU	68	-0.899	12.859	3.795	1.00	1.11
ATOM	1028	Ŋ	HIS	69	-2.420				7.11
ATOM	1029	HN	HIS			11.414	4.454	1.00	0.94
				69	-2.755	10.815	5.155	1.00	1.32
MOTA	1030	CA	HIS	69	-3.114	11.487	3.136	1.00	1.04
ATOM	1031	HA	HIS	69	-2.877	12.437	2.679	1.00	1.25
MOTA	1032	CB	HIS	69	-2.545	10.358	2.243	1.00	1.49
ATOM	1033		HIS	69	-1.750	9.862	2.783		2.12
MOTA	1034		HIS	69				1.00	
ATOM	1035	CG			-2.131	10.798	1.351	1.00	2.27
			HIS	69	-3.570	9.333	1.837	1.00	0.95
ATOM	1036		HIS	69	-3.818	8.195	2.588	1.00	1.43
MOTA	1037		HIS	69	-3.415	7.972	3.453	1.00	1.83
ATOM	1038	CD2	HIS	69	-4.355	9.223	0.717	1.00	1.04
ATOM	1039	HD2	HÍS	69	-4.403	9.946	-0.082	1.00	1.41
MOTA	1040		HIS	69	-4.715	7.452			
				03	-4./13	1.452	1.912	1.00	1.81

» mov	1041	UD1		60	E 003				<b>_</b>
MOTA		HE1		69	-5.097	6.502	2.257	1.00	2.54
MOTA	1042		HIS	69	-5.075	8.032	0.765	1.00	1.53
MOTA	1043		HIS	69	-4.643	11.435	3.341	1.00	1.14
MOTA	1044		HIS	69	-5.392	10.889	2.556	1.00	1.76
MOTA	1045		GLY	70	-5.108	12.065	4.393	1.00	1.49
MOTA	1046		GLY	70	-4.487	12.532	4.990	1.00	1.98
MOTA	1047	CA (	GLY	70	-6.576	12.123	4.665	1.00	1.86
MOTA	1048	HA1 (	GLY	70	-7.071	12.633	3.852	1.00	2.28
ATOM	1049	HA2	GLY	70	-6.746	12.667	5.583	1.00	2.09
MOTA	1050		GLY	70	-7.155	10.716	4.801	1.00	1.81
ATOM	1051		GLY	70	-8.182	10.404	4.232	1.00	2.53
ATOM	1052		ASP	71	-6.513	9.863	5.545		
MOTA	1053		ASP	71				1.00	1.55
MOTA	1054				-5.686			1.00	1.66
			ASP	71	-7.047	8.484	5.701	1.00	1.91
MOTA	1055		ASP	71	-8.126	8.513	5.684	1.00	2.42
MOTA	1056		ASP	71	-6.546	7.620	4.546	1.00	2.67
ATOM	1057	HB1		71	-6.623	6.578	4.813	1.00	3.03
MOTA	1058	HB2		71	-5.514	7.865	4.341	1.00	2.88
MOTA	1059	CG .	ASP	71	-7.397	7.892	3.303	1.00	3.56
ATOM	1060	OD1	ASP	71	-8.476	7.330	3.215	1.00	4.08
MOTA	1061	OD2	ASP	71	-6.960	8.664	2.465	1.00	4.16
ATOM	1062	C .	ASP	71	-6.577	7.889	7.028	1.00	1.46
ATOM	1063		ASP	71	-5.600	8.323	7.605	1.00	1.78
MOTA	1064		PHE	72	-7.260	6.886	7.507	1.00	1.36
ATOM	1065		PHE	72	-8.038	6.546	7.018		1.67
ATOM	1066		PHE	72	-6.849			1.00	
ATOM	1067		PHE			6.248	8.786	1.00	1.48
				72	-6.504	7.007	9.473	1.00	1.75
MOTA	1068		PHE	72	-8.037	5.503	9.399	1.00	2.01
ATOM	1069		PHE	72	-8.374	6.028	10.281	1.00	2.58
MOTA	1070		PHE	72	-7.733	4.503	9.669	1.00	2.43
MOTA	1071		PHE	72	-9.161	5.434	8.395	1.00	2.30
MOTA	1072	CD1	PHE	72	-9.414	4.243	7.704	1.00	2.86
ATOM	1073	HD1	PHE	72	-8.802	3.372	7.887	1.00	3.09
MOTA	1074	CD2	PHE	72	-9.954	6.563	8.158	1.00	2.97
ATOM	1075	HD2	PHE	72	-9.758	7.482	8.691	1.00	3.28
ATOM	1076	CEI		72	-10.459	4.182	6.775	1.00	3.73
ATOM	1077	HE1		72	-10.655	3.264			
ATOM	1078		PHE				6.242	1.00	4.46
				72	-10.999	6.502	7.229	1.00	3.80
MOTA	1079		PHE	72	-11.610	7.374	7.045	1.00	4.54
ATOM	1080		PHE	72	-11.252	5.312	6.537	1.00	4.08
MOTA	1081		PHE	72	-12.058	5.264	5.821	1.00	4.92
MOTA	1082	C	PHE	72	-5.716	5.266	8.500	1.00	1.41
MOTA	1083	0	PHE	72	-5.384	4.430	9.318	1.00	2.20
MOTA	1084	N	TYR	73	-5.120	5.371	7.338	1.00	1.12
ATOM	1085	HN	TYR	73	-5.412	6.059	6.703	1.00	1.48
ATOM	1086		TYR	73	-3.999	4.457	6.972	1.00	1.25
MOTA	1087		TYR	73	-3.774	3.793	7.790	1.00	1.46
ATOM	1088		TYR	73	-4.391	3.635	5.742		
ATOM	1089		TYR	73	-3.531			1.00	1.86
MOTA	1090		TYR	73	-4.726	3.082	5.395	1.00	2.35
ATOM	1091		TYR	-		4.300	4.961	1.00	2.46
ATOM	1092	CD1		73	-5.498	2.670	6.089	1.00	2.08
				73	-5.241	1.585	6.934	1.00	2.58
MOTA	1093		TYR	73	-4.252	1.444	7.347	1.00	2.82
MOTA	1094		TYR	73	-6.779	2.853	5.553	1.00	2.85
MOTA	1095	HD2	TYR	73	-6.978	3.691	4.901	1.00	3.24
MOTA	1096	CE1	TYR	73	-6.264	0.683	7.244	1.00	3.48
MOTA	1097		TYR	73	-6.066	-0.155	7.896	1.00	4.19
MOTA	1098	CE2	TYR	73	-7.802	1.952	5.865	1.00	3.68
MOTA	1099		TYR	73	-8.789	2.093	5.452	1.00	4.49
ATOM	1100		TYR	73	-7.545	0.866	6.710	1.00	3.90
MOTA	1101		TYR	73	-8.554	-0.024	7.013	1.00	5.00
MOTA	1102		TYR	73	-8.689	-0.590	6.249		
ATOM	1103		TYR	73	-2.755			1.00	5.22
ATOM	1104		TYR	73 73		5.273	6.609	1.00	0.95
MOTA					-2.219	5.127	5.529	1.00	1.21
	. 1105	N	PRO	74	-2.273	6.106	7.495	1.00	0.74
MOTA	1106	CA	PRO	74	-1.054	6.895	7.197	1.00	0.82
ATOM	1107	HA	PRO	74	-1.254	7.648	6.453	1.00	1.05
MOTA	1108	CB	PRO	74	-0.746	7.558	8.543	1.00	1.18
MOTA	1109	HB1	PRO	74	-0.786	8.631	8.438	1.00	1.46
MOTA	1110	HB2	PRO	74	0.239	7.261	8.876	1.00	1.28
ATOM	1111	CG	PRO	74	-1.795	7.105	9.566	1.00	1.35
ATOM	1112	HG1		74	-2.229	7.967			1.70
ATOM	1113	HG2		74	-1.330		10.049	1.00	
ATOM	1114	CD	PRO	74		6.468	10.305	1.00	1.61
MOTA	1115	HD2		_	-2.889	6.328	8.828	1.00	1.04
ATOM	1116	HD1		74 74	-3.098	5.393	9.328	1.00	1.24
				74	-3.778	6.929	8.733	1.00	1.14
MOTA	1117	С	PRO	74	0.097	5.988	6.765	1.00	0.65

ATOM	1118	0	PRO	74	0.136	4.822	7.106	1.00	0.66
ATOM	1119	N	PHE	75	1.038	6.503	6.032	1.00	0.56
ATOM	1120	HN	PHE	75	1.000	7.447	5.770		
								1.00	0.61
ATOM	1121	CA	PHE	75	2.179	5.651	5.605	1.00	0.45
ATOM	1122	HA	PHE	· 75	1.816	4.659	5.360	1.00	0.48
ATOM	1123	CB	PHE	75	2.859	6.266	4.379	1.00	0.42
MOTA	1124		PHE	75	3.761	5.718	4.153	1.00	0.44
MOTA	1125	HB2	PHE	75	3.104	7.298	4.582	1.00	0.45
MOTA	1126	CG	PHE	75	1.915	6.190	3.200	1.00	0.48
MOTA	1127	CD1		75	1.764	4.986	2.501	1.00	0.41
MOTA	1128	HD1	PHE	75	2.329	4.115	2.797	1.00	0.45
ATOM	1129	CD2	PHE	75	1.184	7.320	2.812	1.00	0.74
ATOM	1130		PHE	75	1.300	8.249			
							3.349	1.00	0.90
MOTA	1131	CEL		75	0.882	4.911	1.415	1.00	0.50
MOTA	1132	HE1	PHE	75	0.767	3.982	0.877	1.00	0.53
ATOM	1133	CE2	PHE	75	0.304	7.245	1.724	1.00	0.85
MOTA	1134	HE2	PHE	75	-0.258	8.117	1.423	1.00	1.09
ATOM	1135	CZ	PHE	75	0.154	6.041	1.026	1.00	0.69
· ATOM	1136	HZ	PHE	75	-0.526	5.983	0.188	1.00	0.80
ATOM	1137	C	PHE	75		5.561			
					3.159		6.776	1.00	0.43
MOTA	1138	0	PHE	75	3.111	6.360	7.690	1.00	0.50
ATOM	1139	N	ASP	76	4.020	4.582	6.782	1.00	0.37
ATOM	1140	HN	ASP	76	4.028	3.929	6.050	1.00	0.32
MOTA	1141	CA	ASP	76	4.967	4.432	7.927	1.00	0.43
MOTA	1142	HA	ASP	76	4.551	4.906	8.804	1.00	0.50
MOTA	1143	CB	ASP	76	5.180	2.946	8.215	1.00	0.46
ATOM	1144								
			ASP	76	4.224	2.467	8.365	1.00	0.49
ATOM	1145	HB2	ASP	76	5.784	2.834	9.104	1.00	0.54
ATOM	1146	CG	ASP	76	5.892	2.295	7.028	1.00	0.38
MOTA	1147	OD1	ASP	76	6.468				
						1.236	7.218	1.00	0.45
MOTA	1148	OD2	ASP	76	5.846	2.864	5.950	1.00	0.30
ATOM	1149	C	ASP	76	6.314	5.074	7.596	1.00	0.42
MOTA	1150	0	ASP	76	7.314	4.770	8.216	1.00	0.54
ATOM									
	1151	N	GLY	77	6.347	5.958	6.632	1.00	0.35
ATOM	1152	HN	GLY	77	5.525	6.187	6.151	1.00	0.36
ATOM	1153	CA	GLY	77	7.634	6.625	6.267	1.00	0.38
MOTA	1154	HA1		77					
					8.378	6.388	7.004	1.00	0.45
MOTA	1155	HA2	GLY	77	7.484	7.696	6.238	1.00	0.44
MOTA	1156	С	GLY	<i>77</i>	8.084	6.131	4.884	1.00	0.31
ATOM	1157	ō	GLY	77					
	1157				7.262	5.767	4.068	1.00	0.37
MOTA	1158	N	PRO	78	9.370	6.117	4.603	1.00	0.33
MOTA	1159	CA	PRO	78	9.856	5.651	3.274	1.00	0.36
MOTA	1160	HA	PRO	78	9.435	6.254	2.488	1.00	0.42
ATOM	1161	CB	PRO	78	11.364	5.903	3.359	1.00	0.46
ATOM	1162	HB1	PRO	78	11.671	6.542	2.545	1.00	0.56
MOTA	1163	HB2	PRO	78	11.892	4:962	3.303	1.00	0.48
ATOM	1164	CG			11.675				
			PRO	78	11.0/5	6.592	4.694	1.00	0.64
MOTA	1165	HG1	PRO	78	11.965	7.616	4.516	1.00	0.87
MOTA	1166	HG2	PRO	78	12.478	6.068	5.194	1.00	0.83
ATOM	1167	CD	PRO	78	10.418	6.562	5.563	1.00	_
									0.45
ATOM	1168		PRO	78	10.535	5.848	6.369	1.00	0.48
ATOM	1169	HD1	PRO	78	10.187	7.544	5.944	1.00	0.49
MOTA	1170	C	PRO	78	9.564	4.165	3.027	1.00	0.30
ATOM	1171	ō		78					
			PRO		8.860	3.808	2.105	1.00	0.28
MOTA	1172	N	SER	79	10.102	3.297	3.840	1.00	0.31
MOTA	1173	HN	SER	79	10.670	3.604	4.577	1.00	0.35
ATOM	1174	CA	SER	79	9.855	1.837	3.647	1.00	0.30
ATOM	1175	HA	SER	79	9.916	1.599			
							2.595	1.00	0.30
MOTA	1176	CB	SER	79	10.911	1.037	4.410	1.00	0.37
ATOM	1177	HB1	SER	79	11.888	1.465	4.225	1.00	0.42
MOTA	1178	HB2		79	10.901	0.013	4.076		0.39
ATOM	1179							1.00	
		OG	SER	79	10.617	1.080	5.800	1.00	0.38
ATOM	1180	HG	SER	79	11.173	1,752	6.201	1.00	0.98
MOTA	1181	C	SER	79	8.463	1,470	4.173	1.00	0.27
ATOM	1182				7 000		7.1/3	1.00	
		0	SER	79	7.888	2.183	4.971	1.00	0.25
MOTA	1183	N	GLY	80	7.927	0.356	3.734	1.00	0,31
ATOM	1184	HN	GLY	80	8.420	-0.200	3.095	1.00	0.37
MOTA	1185	CA	GLY	80	£ F36				
					6.576	-0.081	4.207	1.00	0.30
MOTA	1186	HA1		80	6.224	0.586	4.977	1.00	0.31
MOTA	1187	HA2	GLY	80	6.646	-1.083	4.607	1.00	0.36
ATOM	1188	c	GLY	80					
	1100				5.584	-0.070	3.042	1.00	0.25
ATOM	1189	0	GLY	80	5.850	-0.601	1.981	1.00	0.25
MOTA	1190	N	LEU	81	4.440	0.531	3.232	1.00	0.23
ATOM	1191	HN	LEU	81					
MOTA	1192		LEU		4.246	0.951	4.096	1.00	0.25
		CA		81	3.428	0.577	2.138	1.00	0.21
ATOM	1193	HA	LEU	81	3.259	-0.417	1.761	1.00	0.22
MOTA	1194	CB	LEU	81	2.123	1.164	2.692	1.00	0.24
					~ . 14.	~ . 704	2.032	1.44	u.z.

ATOM	1195	HB1 L	.PII	81	1.587	1.658	1.896	1.00	0.25
MOTA	1196	HB2 L		81	2.356	1.881	3.465	1.00	0.29
MOTA	1197		ĒÜ	81					
					1.240	0.058	3.283	1.00	0.28
MOTA	1198		ΣŲ	81	1.856	-0.678	3.779	1.00	0.31
MOTA	1199	CD1 L	ŒU	81	0.265	0.680	4.285	1.00	0.33
MOTA	1200	HD11 L	ŒU	81	0.071	1.706	4.009	1.00	1.05
ATOM	1201	HD12 L	EU	81	0.696	0.649	5.274	1.00	1.10
ATOM			EU	81	-0.662				
						0.125	4.278	1.00	1.06
MOTA	1203	CD2 L		81	0.426	-0.606	2.168	1.00	0.31
MOTA		HD21 L		81	1.087	-0.997	1.412	1.00	1.02
MOTA	1205	HD22 L	ŒU	81	-0.233	0.126	1.724	1.00	1.09
ATOM	1206	HD23 L	ŒU	81	-0.161	-1.411	2.584	1.00	1.06
ATOM	1207		EU	81	3.953	1.475	1.017	1.00	0.20
ATOM	1208	-	EU	81	3.988	2.679	1.141	1.00	
	1209								0.22
ATOM			EU	82	4.366	0.899	-0.078	1.00	0.18
MOTA	1210		LEU	82	4.334	-0.077	-0.162	1.00	0.18
MOTA	1211	ÇA I	LEU	82	4.901	1.728	-1.195	1.00	0.18
ATOM	1212	HA I	LEU	82	5.519	2.520	-0.799	1.00	0.19
ATOM	1213	CB I	LEU	82	5.728	0.840	-2.128	1.00	0.18
ATOM	1214	HB1 I		82	6.235	1.457	-2.854	1.00	0.20
MOTA	1215	HB2 I		82					
					5.071	0.151	-2.640	1.00	0.20
ATOM	1216		LEU	82	6.763	0.050	-1.323	1.00	0.18
MOTA	1217		LEU	82	6.262	-0.523	-0.556	1.00	0.22
MOTA	1218	CD1 I		82	7.513	-0.898	-2.259	1.00	0.17
ATOM	1219	HD11 I	JEU	82	8.102	-0.321	-2.957	1.00	0.97
MOTA	1220	HD12 I	LEU	82	6.802	-1.503	-2.802	1.00	0.95
ATOM	1221	HD13 I	us	82	8.163	-1.537	-1.681	1.00	
ATOM	1222	CD2		. 82					0.98
					7.764	1.010	-0.675	1.00	0.23
MOTA	1223	HD21 I		82	8.019	1.790	-1.375	1.00	1.03
ATOM	1224		LEU	82	8.657	0.466	-0.403	1.00	1.07
MOTA	1225	HD23 I	LEU	82	7.326	1.447	0.209	1.00	1.02
ATOM	1226	C I	LEU	82	3.740	2.329	-1.986	1.00	0.19
MOTA	1227	0 1	LEU	82	3.882	3.341	-2.646	1.00	0.21
ATOM	1228		ALA	83	2.594	1.711	-1.919		0.21
MOTA	1229		ALA	83				1.00	
					2.512	0.899	-1.376	1.00	0.24
MOTA	1230		ALA	83	1.410	2.225	-2.662	1.00	0.22
MOTA	1231		ALA	83	1.217	3.251	-2.381	1.00	0.22
ATOM	1232	CB 1	ALA	83	1,668	2.140	-4.171	1.00	0.23
ATOM	1233	HB1 2	ALA	83	2,522	2.746	-4.429	1.00	0.98
ATOM	1234		ALA	83	0.801	2.497	-4.705	1.00	1.00
ATOM	1235		ALA	83	1.860	1.113			
ATOM	1236						-4.445	1.00	1.05
			ALA	83	0.204	1.350	-2.317	1.00	0.27
MOTA	1237		ALA	83	0.342	0.301	-1.720	1.00	0.36
ATOM	1238		HIS	84	-0.976	1.762	-2.686	1.00	0.24
MOTA	1239	HN I	HIS	84	-1.075	2.609	-3.170	1.00	0.20
MOTA	1240	CA I	HIS	84	-2.173	0.933	-2.370	1.00	0.30
MOTA	1241		HIS	84	-1.940	-0,108	-2.542	1.00	0.36
ATOM	1242		HIS	84	-2.562	1.127			
ATOM	1243		HIS	84			-0.903	1.00	0.40
					-1.695	0.965	-0.278	1.00	0.48
MOTA	1244		HIS	84	-3.332	0.419	-0.638	1.00	0.45
ATOM	1245		HIS	84	-3.074	2.525	-0.692	1.00	0.44
MOTA	1246	ND1	HIS	84	-4.384	2.781	-0.321	1.00	1.32
ATOM	1247	HD1	HIS	84	-5.084	2.112	-0.169	1.00	2.02
ATOM	1248	CD2		84	-2.465	3.752	-0.788	1.00	0.74
ATOM	1249	HD2 1		84	-1.432	3.915	-1.060	1.00	1.58
MOTA	1250	CE1		84	-4.521	4.114	-0.208		
ATOM	1251	HE1		84				1.00	1.21
ATOM	1252		HIS		-5.441	4.606	0.071	1.00	1.87
ATOM				84	-3.381	4.754	-0.482	1.00	0.53
-	1253		HIS	84	-3.337	1.343	-3.274	1.00	0.25
ATOM	1254	0 1	HIS	84	-3.347	2.417	-3.843	1.00	0.23
MOTA	1255	N 3	ALA	85	-4.313	0.489	-3.417	1.00	0.27
MOTA	1256	HN .	ALA	85	-4.279	-0.374	-2.954	1.00	0.34
ATOM	1257		ALA	85	-5.474	0.817	-4.291	1.00	
ATOM	1258		ALA	85	-5.582				0.24
ATOM						1.890	-4.364	1.00	0.22
	1259		ALA	85	-5.236	0.231	-5.685	1.00	0.25
MOTA	1260		ALA	85	-5.079	-0.835	-5.605	1.00	1.05
MOTA	1261		ALA	85	-4.364	0.690	-6.126	1.00	1.05
ATOM	1262	нвз 2	ALA	85	-6.097	0.420	-6.308	1.00	1.06
ATOM	1263		ALA	85	-6.748	0.210	-3.698	1.00	0.26
MOTA	1264		ALA	85	-6.694				
ATOM	1265					-0.611	-2.804	1.00	0.33
			PHE	86	-7.892	0.605	-4.198	1.00	0.28
MOTA	1266		PHE	86	-7.905	1.264	-4.922	1.00	0.31
MOTA	1267		PHE	86	-9.179	0.053	-3.677	1.00	0.34
MOTA	1268		PHE	86	-9.000	-0.443	-2.737	1.00	0.39
MOTA	1269	CB :	PHE	86	-10.170	1.205	-3.471	1.00	0.36
ATOM	1270		PHE	86	-11.177	0.821	-3.459	1.00	0.42
MOTA	1271	HB2		86	-10.068	1.913			
			ad	-	-10.000	T.313	-4.279	1.00	0.33

MOTA	1272	CG	PHE	86	-9.877	1.896	-2.159	1.00	0.39
MOTA	1273	CD1		86	-8.784	2.764	-2.050	1.00	0.46
ATOM	1274	HD1		86	-8.146	2.939	-2.903		
								1.00	0.67
MOTA	1275	CD2		86	-10.703	1.670	-1.051	1.00	0.67
MOTA	1276		PHE	86	-11.546	1.001	-1.133	1.00	0.91
ATOM	1277	CE1	PHE	86	-8.516	3.406	-0.835	1.00	0.50
MOTA	1278	HE1	PHE	86	-7.673	4.075	-0.751	1.00	0.69
ATOM	1279		PHE	86	-10.435	2.311	0.165	1.00	0.74
ATOM	1280		PHE	86	-11.071	2.136	1.020	1.00	
									1.02
MOTA	1281		PHE	86	-9.342	3.179	0.273	1.00	0.54
MOTA	1282		PHE	86	-9.135	3.674	1.211	1.00	0.62
ATOM	1283	С	PHE	86	-9.746	-0.940	-4.710	1.00	0.36
MOTA	1284	0	PHE	86	-9.480	-0.812	-5.889	1.00	0.34
MOTA	1285		PRO	87	-10.516	-1.926	-4.293	1.00	0.43
MOTA	1286		PRO	87	-11.082	-2.914	-5.257	1.00	
									0.46
MOTA	1287		PRO	87	-10.296	-3.524	-5.665	1.00	0.53
ATOM .	1288	CB	PRO	87	-11.990	-3.770	-4.370	1.00	0.60
MOTA	1289	HB1	PRO	87	-11.644	-4.792	-4.377	1.00	0.69
MOTA	1290	HB2	PRO	87	-13.004	-3.727	-4.742	1.00	0.73
ATOM	1291	CG	PRO	87	-11.943	-3,225	-2.937	1.00	0.58
ATOM	1292		PRO	87	-11.694	-4.022	-2.253	1.00	0.61
MOTA	1293	HG2	PRO	87	-12.905	-2.808	-2.676	1.00	0.66
MOTA	1294	CD	PRO	87	-10.872	-2.135	-2.861	1.00	0.50
MOTA	1295		PRO	87	-11.277	-1.235	-2.421	1.00	0.50
ATOM	1296	HD1	PRO	87	-10.014	-2.484	-2.309	1.00	0.52
MOTA	1297	С	PRO	87	-11.895	-2.246	-6.379	1.00	0.40
MOTA	1298	0	PRO	87	-12.221	-1.078	-6.299	1.00	0.42
MOTA	1299	N	PRO	88	-12.221	-2.981	-7.419	1.00	
					-12.221				0.44
MOTA	1300	CA	PRO	88	-13.007	-2.416	-8.554	1.00	0.48
MOTA	1301	HA	PRO	88	-12.443	-1.645	-9.053	1.00	0.52
MOTA	1302	CB	PRO	88	-13.163	-3.622	-9.488	1.00	0.61
ATOM	1303	HB1	PRO	88	-12.604	-3.449	-10.395	1.00	0.83
MOTA	1304	HB2	PRO	88	-14.204	-3.772	-9.728	1.00	0.74
ATOM	1305	CG	PRO	88	-12.609	-4.863	-8.781	1.00	0.57
ATOM	1306								
			PRO	88	-11.945	-5.395	-9.446	1.00	0.71
ATOM	1307		PRO	88	-13.425	-5.508	-8.488	1.00	0.64
MOTA	1308	CD	PRO	88	-11.835	-4.413	-7.540	1.00	0.56
ATOM	1309	HD2	PRO	88	-12.146	-4.977	-6.671	1.00	0.62
MOTA	1310	HD1	PRO	88	-10.773	-4.503	-7.702	1.00	0.65
ATOM	1311	C	PRO	88	-14.372	-1.873	-8.109	1.00	0.47
ATOM	1312	ŏ	PRO	88					
					-15.380	-2.551	-8.172	1.00	0.88
MOTA	1313	N	GLY	89	-14.400	-0.647	-7.661	1.00	0.63
MOTA	1314	HN	GLY	89	-13.571	-0.129	-7.626	1.00	1.01
ATOM	1315	CA	GLY	89	-15.681	-0.026	-7.209	1.00	0.65
ATOM	1316	HA1	GLY	89	-15.536	0.422	-6.239	1.00	0.62
ATOM	1317	HA2	GLY	89	-16.455	-0.778	-7.148	1.00	0.78
ATOM	1318	c	GLY	89	-16.092	1.057			
ATOM	1319	ŏ					-8.210	1.00	0.74
			GLY	89	-15.541	1.151	~9.289	1.00	0.84
MOTA	1320	N	PRO	90	-17.044	1.878	-7.852	1.00	0.95
MOTA	1321	CA	PRO	90	-17.499	2.973	-8.750	1.00	1.19
MOTA	1322	HA	PRO	90	-17.819	2.565	-9.697	1.00	1.37
ATOM	1323	CB	PRO	90	-18.720	3.532	-7.990	1.00	1.55
ATOM	1324		PRO	90	-19.602	3.432	-8.605	1.00	1.85
ATOM	1325		PRO	90	-18.572	4.567	-7.740	1.00	1.74
ATOM	1326	CG	PRO	90	-18.913	2.724			
							-6.702	1.00	1.46
MOTA	1327		PRO	90	-19.828	2.155	-6.763	1.00	1.60
MOTA	1328		PRO	90	-18.959	3.396	-5.857	1.00	1.57
MOTA	1329	CD	PRO	90	-17.729	1.769	-6.539	1.00	1.17
MOTA	1330	HD2	PRO	90	-17.083	2.099	-5.736	1.00	1.17
MOTA	1331	HD1	PRO	90	-18.067	0.759	-6.375	1.00	1.28
MOTA	1332	C	PRO	90	-16.375	4:011	-8.972	1.00	1.14
MOTA	1333	ō	PRO	90	-15.269	3.649	-9.320	1.00	1.53
MOTA	1334								
		N	ASN	91	-16.624	5,282	-8.790	1.00	1.17
MOTA	1335	HN	ASN	91	-17.514	5.578	-8.517	1.00	1.40
MOTA	1336	CA	asn	91	-15.541	6.286	-9.008	1.00	1.38
MOTA	1337	HA	asn	91	-15.147	6.169	-10.005	1.00	1.58
MOTA	1338	CB	ASN	91	-16.116	7:700	-8.857	1.00	1.87
ATOM	1339		ASN	91	-15.336	8.372	-8.532	1.00	2.33
MOTA	1340		ASN	91	-16.908				
ATOM	1341					7.686	-8.122	1.00	1.96
		CG	ASN	91	-16.678		-10.197	1.00	2.69
MOTA	1342		ASN	91	-16.132	7.890	-11.242	1.00	3.20
MOTA	1343		asn	· 91	-17.748	8.931	-10.212	1.00	3.47
MOTA	1344	HD21		91	-18.186	9.176	-9.370	1.00	3.59
ATOM	1345	HD22	ASN	91	-18.112		-11.064	1.00	4.20
ATOM	1346	C	ASN	91	-14.404	6.098	-7.992	1.00	1.15
ATOM	1347	ŏ	ASN	91					
MOTA	1348	Ŋ			-13.242	6.135	-8.344	1.00	1.26
ALUM	7340	7.4	TYR	92	-14 719	5 924	_£ 735	1 00	דח ר

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ATOM	1349	HN	TYR	92	-15.660	5.916	-6.462	1 00	1 00
ATOM	1350	CA	TYR	92				1.00	1.08
					-13.639	5.768	-5.711	1.00	0.97
ATOM	1351	HA	TYR	92	-12.994	6.632	-5.739	1.00	1.14
MOTA	1352	CB	TYR	92	-14.262	5.652	-4.319	1.00	1.09
ATOM	1353	HB1	TYR	92	-13.543	5.214	-3.643	1.00	1.62
ATOM	1354	HB2	TYR	92	-15.135	5.020	-4.369	1.00	1.45
ATOM .	1355	CG	TYR	92	-14.656	7.018	-3.810	1.00	1.52
ATOM	1356		TYR	92	-13.672	7.979	-3.549		
ATOM	1357	_	TYR	92				1.00	2.14
					-12.631	7.747	-3.719	1.00	2.46
MOTA	1358	CD2	TYR	92	-16.006	7.320	-3.588	1.00	2.44
ATOM	1359	HD2	TYR	. 92	~16.766	6.580	-3.789	1.00	2.86
MOTA	1360	CEl	TYR	92	-14.037	9.241	-3.066	1.00	3.06
MOTA	1361	HE1	TYR	92	-13.278	9.982	-2.865	1.00	3.78
ATOM	1362	CE2	TYR	92	-16.370	8.582	-3.107	1.00	3.33
ATOM	1363	HE2	TYR	92	-17.411	8.815			
ATOM	1364	CZ	TYR	92			-2.936	1.00	4.19
MOTA	1365				-15.386	9.542	-2.846	1.00	3.50
		ОН	TYR	92	-15.746	10.786	-2.368	1.00	4.57
MOTA	1366	HH	TYR	92	-15.602	10.791	-1.419	1.00	4.91
ATOM	1367	С	TYR	92	~12.808	4.508	-5.966	1.00	0.78
MOTA	1368	0	TYR	92	-11.605	4.506	-5.798	1.00	0.81
MOTA	1369	N	GLY	93	-13.436	3.430	-6.337	1.00	0.64
MOTA	1370	HN	GLY	93	-14.410	3.441	-6.445	1.00	0.70
ATOM	1371	CA	GLY	93	-12.674	2.170	-6.560	1.00	
ATOM	1372		GLY	93	-13.366				0.51
ATOM	1373					1.366	-6.740	1.00	0.51
		HA2		93	-12.090	1.947	-5.67B	1.00	0.51
ATOM	1374	С	GLY	93	-11.739	2.310	-7.761	1.00	0.49
ATOM	1375	0	GLY	93	-11.832	3.242	-8.534	1.00	0.61
ATOM	1376	N	GLY	94	-10.844	1.373	-7.923	1.00	0.45
ATOM	1377	HN	GLY	94	-10.799	0.627	-7.288	1.00	0.44
MOTA	1378	CA	GLY	94	-9.902	1.420	-9.075	1.00	
ATOM	1379		GLY	94	-10.459				0.55.
ATOM	1380		GLY			1.569	-9.988	1.00	0.63
				94	-9.363	0.485	-9.133	1.00	0.58
ATOM	1381	C	GLY	94	-8.905	2.569	-8.901	1.00	0.60
MOTA.	1382	0	GLY	94	-8.109	2.838	-9.772	1.00	1.14
MOTA	1383	N	ASP	95	-8.933	3.252	-7.790	1.00	0.24
MOTA	1384	HN	ASP	95	-9.581	3.028	-7.089	1.00	0.52
ATOM	1385	CA	ASP	95	-7.976	4.382	-7.597	1.00	0.24
ATOM	1386	HA	ASP	95	-7.888	4.939	-8.518		
ATOM	1387	CB	ASP	95				1.00	0.28
ATOM					-8.493	5.303	-6.491	1.00	0.26
	1388		ASP	95	-9.500	5.617	-6.724	1.00	0.28
MOTA	1389		ASP	95	-7.853	6.170	-6.415	1.00	0.30
MOTA	1390	CG	ASP	95	-8.494	4.549	-5.162	1.00	0.28
MOTA	1391	QD1	ASP	95	-8.543	5.200	-4.132	1.00	1.08
MOTA	1392	OD2	ASP	95	-8.440	3.331	-5.198	1.00	1.14
MOTA	1393	С	ASP	95	-6.605	3.827	-7.202	1.00	0.23
ATOM	1394	ŏ	ASP	95	-6.479				
MOTA	1395	N	ALA			2.683	-6.815	1.00	0.24
				96	-5.573	4.626	-7.297	1.00	0.23
MOTA	1396	HN	ALA	96	-5.692	5.546	-7.614	1.00	0.23
MOTA	1397	CA	ALA	96	-4.215	4.131	-6.926	1.00	0.25
MOTA	1398	HA	ALA	96	-4.307	3:360	-6.175	1.00	0.25
MOTA	1399	CB	ALA	96	-3.527	3.553	-8.164	1.00	0.30
MOTA	1400	HB1	ALA	96	-2.528	3.236	-7.905	1.00	1.08
MOTA	1401	HB2	ALA	96	-3.476	4.309	-8.934	1.00	1.08
ATOM	1402		ALA	96	-4.090	2.706	-8.528	1.00	
MOTA	1403	C	ALA	96	-3.375				1.03
ATOM	1404	ŏ	ALA	96		5.284	-6.372	1.00	0.25
ATOM	1405				-3.222	6.313	-7.005	1.00	0.29
		N	HXS	97	-2.831	5.113	-5.192	1.00	0.25
MOTA	1406	HN	HXS	97	-2.976	4,271	-4.710	1.00	0.28
MOTA	1407	CA	HXS	97	-1.996	6.187	-4.574	1.00	0.27
MOTA	1408	HA	HXS	97	-2.010	7.068	-5.198	1.00	0.28
MOTA	1409	CB	HXS	97	-2.564	6.537	-3.197	1.00	0.33
MOTA	1410	HB1	HXS	97	-1.969	7.319	-2.750	1.00	
ATOM	1411		HXS	97	-2.540				0.44
ATOM	1412	CG	HXS	97		5.661	-2.566	1.00	0.39
ATOM	1413		HXS		-3.983	7.009	-3.349	1.00	0.37
				97	-4.697	7.052	-2.163	1.00	0.80
MOTA	1414		HXS	97	-4.783	7.420	-4.384	1.00	0.55
MOTA	1415		HXS	97	-4.517	7.497	-5.428	1.00	0.94
MOTA	1416	CE1	HXS	97	-5.918	7.487	-2.498	1.00	0.86
MOTA	1417		HXS	97	-6.724	7.632	-1.795	1.00	1,24
ATOM	1418		HXS	97	-6.018	7.722	-3.819		
ATOM	1419		HXS	97	-6.812			1.00	0.59
ATOM	1420	C	HXS	97		8.044	-4.294	1.00	0.72
ATOM	1421				-0.552	5.700	-4.420	1.00	0.26
		0	HXS	97	-0.299	4.525	-4.237	1.00	0.39
MOTA	1422	N	PHE	98	0.391	6.604	-4.496	1.00	0.18
ATOM	1423	HN	PHE	98	0.147	7.540	-4.648	1.00	0.23
MOTA	1424	CA	PHE	98	1.832	6.230	-4.360	1.00	0.17
MOTA	1425	HA	PHE	98	1.921	5.190	-4.085	1.00	0.18
								00	U. 10

MOTA	1426	CB	PHE	98	2.543	6.472	-5.691	1.00	0.18
MOTA	1427	HB1		98	3.611	6.464	-5.536	1.00	0.21
MOTA	1428		PHE	98	2.243	7.431	-6.085	1.00	0.20
ATOM	1429	CG	PHE	98	2.169	5.391	-6.674	1.00	0.19
MOTA	1430	CD1	PHE	98	3.114	4.428	-7.048	1.00	0.22
MOTA	1431	HD1	PHE	98	4.110	4.456	-6.631	1.00	0.25
ATOM	1432		PHE	98	0.880	5.355	-7.214	1.00	0.22
ATOM	1433	HD2		98	0.151	6.098	-6.924	1.00	0.24
ATOM	1434	CE1		98	2.768	3.429			
ATOM	1435		PHE	98		2.685	-7.963	1.00	0.25
ATOM	1436	CE2			3.496	2.085	-8.252	1.00	0.29
			PHE	98	0.533	4.355	-8.127	1.00	0.26
MOTA	1437	HE2	PHE	98	-0.462	4.327	-8.542	1.00	0.31
MOTA	1438	CZ	PHE	98	1.478	3.392	-8.503	1.00	0.26
ATOM	1439	HZ	PHE	98	1.214	2.622	-9.211	1.00	0.30
MOTA	1440	С	PHE	98	2.487	7.104	-3.286	1.00	0.17
MOTA	1441	0	PHE	98	2.081	8.226	~3.058	1.00	0.19
ATOM	1442	N	ASP	99	3.498	6.604	-2.625	1.00	0.19
ATOM	1443	HN	ASP	99	3.813	5.693	-2.820	1.00	0.22
ATOM	1444	CA	ASP	99	4.167	7.424	-1.570	1.00	0.20
ATOM	1445	HA	ASP	99	3.421	7.956	-0.998	1.00	0.20
ATOM	1446	CB	ASP	99	4.973	6.516	-0.638	1.00	0.25
ATOM	1447	HB1		99	5.567	7.122	0.029	1.00	
ATOM	1448	HB2		99	5.624				0.28
ATOM	1449	CG	ASP	99		5.884	-1.226	1.00	0.30
ATOM	1450				4.023	5.646	0.180	1.00	0.41
-		OD1	ASP	99	2.838	5.680	-0.100	1.00	0.89
ATOM	1451	OD2		99	4.497	4.968	1.079	1.00	0.27
MOTA	1452	Ċ	ASP	99	5.123	8.426	-2.224	1.00	0.21
MOTA	1453	0	ASP	99	6.020	8.054	-2.954	1.00	0.25
MOTA	1454	N	ASP	100	4.946	9.694	-1.962	1.00	0.23
MOTA	1455	HN	ASP	100	4.222	9.976	-1.365	1.00	0.23
MOTA	1456	CA	ASP	100	5.857	10.710	-2.565	1.00	0.29
ATOM	1457	HA	ASP	100	6.169	10.379	-3.545	1.00	0.31
ATOM	1458	CB	ASP	100	5.127	12.049	-2.684	1.00	0.34
ATOM	1459	HB1		100	5.130	12.544	-1.727		
ATOM	1460	HB2		100	4.109			1.00	0.34
ATOM	1461	CG	ASP			11.879	-2.999	1.00	0.34
				100	5.844	12.929	-3.710	1.00	0.43
MOTA	1462	OD1		100	5.240	13.887	-4.164	1.00	1.21
ATOM	1463	OD2		100	6.984	12.630	-4.025	1.00	1.12
MOTA	1464	Ç	ASP	100	7.085	10.885	-1.667	1.00	0.30
MOTA	1465	0	ASP	100	8.032	11.559	-2.018	1.00	0.32
MOTA	1466	N	ASP	101	7.074	10.280	-0.510	1.00	0.31
MOTA	1467	HN	ASP	101	6.298	9.741	-0.249	1.00	0.32
ATOM	1468	CA	ASP	101	8.236	10.407	0.415	1.00	0.33
ATOM	1469	HA	ASP	101	8.647	11:403	0.345	1.00	0.36
ATOM	1470	CB	ASP	101	7.778	10.142	1.851		
ATOM	1471	HB1		101	8.641			1.00	0.39
ATOM	1472		ASP	101		10.060	2.495	1.00	0.41
ATOM	1473	CG	ASP		7.216	9.220	1.884	1.00	0.39
ATOM	1474			101	6.896	11.296	2.330	1.00	0.45
		OD1	ASP	101	7.027	12.380	1.786	1.00	1.25
MOTA	1475		ASP	101	6.104	11.076	3.231	1.00	1.09
MOTA	1476	Ç	ASP	101	9.304	9.385	0.028	1.00	0.30
MOTA	1477	0	ASP	101	10.411	9.405	0.529		0.29
ATOM	1478	N	GLU	102	8.971	8.484	-0.849	1.00	0.30
MOTA	1479	HN	GLU	102	8.068	8.484	-1.230	1.00	0.31
MOTA	1480	CA	GLU	102	9.950	7.444	-1.266	1.00	0.29
MOTA	1481	HA	GLU	102	10.649	7.263	-0.463	1.00	0.30
MOTA	1482	CB	GLU	102	9.195	6.155	-1.585	1.00	0.35
MOTA	1483	HB1	GLU	102	9.873	5.437	-2.020	1.00	0.36
ATOM	1484		GLU	102	8.397	6.368	-2.282	1.00	0.40
ATOM	1485	CG	GLU	102	8.611	5.584	-0.293	1.00	0.46
ATOM	1486		GLU	102	8.020	6.342	0.200		1.18
ATOM	1487		GLU	102	9.415			1.00	
ATOM	1488	CD	GLU	102		5.276	0.356	1.00	1.03
ATOM	1489				7.724	4.381	-0.616	1.00	0.83
ATOM			GLU	102	7.601	4.060	-1.786	1.00	1.63
	1490	OE2	GLU	102	7.184	3.801	0.314	1.00	0.87
MOTA	1491	C	GLU	102	10.707	7.917	-2.508	1.00	0.25
MOTA	1492	0	GLU	102	10.359	8.910	-3.115	1.00	0.25
MOTA	1493	N	THR	103	11.741	7.213	-2.886	1.00	0.25
MOTA	1494	HN	THR	103	12.003	6.416	-2.379	1.00	0.28
MOTA	1495	CA	THR	103	12.525	7.620	-4.088	1.00	0.23
ATOM	1496	HA	THR	103	12.356	8.665	-4.301	1.00	0.23
MOTA	1497	CB	THR	103	14.016	7.383	-3.824	1.00	0.27
MOTA	1498	HB	THR	103	14.169	6.359	-3.521	1.00	0.30
MOTA	1499	OG1	THR	103	14.455				
ATOM	1500	HG1	THR	103	15.334	8.252	-2.789	1.00	0.29
ATOM	1501	CG2	THR	103		8.564	-3.016	1.00	0.86
ATOM		HG21		2 0 -	14.820	7.656 7.777	-5.098	1.00	0.29
			****	103 .	15.864	1 777	-A BAR	1 00	. 110

ATOM		HG22		103	14.457	8.557	-5.569	1.00	1.08
MOTA	1504	HG23	THR	103	14.710	6.824	-5.779	1.00	1.01
ATOM	1505	С	THR	103	12.083	6.777	-5.281	1.00	
ATOM	1506	Ö	THR	103	12.417	5.614	-5.394		0.22
MOTA	1507	Ň	TRP	104	11.332			1.00	0.23
ATOM	1508	HN	TRP	104		7.358	-6.175	1.00	0.21
ATOM	1509				11.076	8.297	-6.063	1.00	0.23
		CA	TRP	104	10.867	6.598	-7.364	1.00	0.21
ATOM	1510	HA	TRP	104	10.750	5.556	-7.104	1.00	0.20
ATOM	1511	CB	TRP	104	9.525	7.165	-7.831	1.00	0.23
MOTA	1512	HB1		104	9.188	6.623	-8.702	1.00	0.24
ATOM	1513	HB2	TRP	104	9.641	8.210	-8.078	1.00	0.25
ATOM	1514	CG	TRP.	104	8.520	7.018	-6.731	1.00	0.24
MOTA	1515	CD1	TRP	104	8.098	8.019	-5.924	1.00	0.31
ATOM	1516	HD1	TRP	104	8.427	9.045	-5.972	1.00	0.36
MOTA	1517	CD2	TRP	104	7.811	5.821	-6.300		
ATOM	1518	NE1		104	7.176		-5.026	1.00	0.21
ATOM	1519	HE1		104	6.718	7.512		1.00	0.31
ATOM	1520	CE2		104		8.030	-4.331	1.00	0.36
ATOM	1521	CE3			6.963	6.162	-5.220	1.00	0.24
			TRP	104	7.819	4.486	-6.739	1.00	0.18
MOTA	1522	HE3		104	8.458	4.198	-7.559	1.00	0.19
ATOM	1523	CZ2		104	6.153	5.213	-4.596	1.00	0.23
MOTA	1524	HZ2		104	5.515	5.499	-3.774	1.00	0.27
MOTA	1525	CZ3	TRP	104	7.005	3.527	-6.114	1.00	0.20
MOTA	1526	HZ3	TRP	104	7.019	2.504	-6.460	1.00	0.23
ATOM	1527	CH2	TRP	104	6.173	3.891	-5.045	1.00	0.21
ATOM	1528	HH2	TRP	104	5.548	3.150	-4.568	1.00	
MOTA	1529	C	TRP	104	11.911	6.732			0.23
MOTA	1530	ō	TRP	104	12.276		-8.474	1.00	0.21
ATOM	1531	Ň	THR	105		7.824	-8.864	1.00	0.24
ATOM	1532	HN			12.403	5.630	-8.973	1.00	0.20
ATOM			THR	105	12.098	4.763	-8.633	1.00	0.19
	1533	CA	THR	105	13.437	5.685		1.00	0.21
MOTA	1534	HA	THR	105	13.415	6.652	-10.525	1.00	0.24
MOTA	1535	CB	THR	105	14.817	5.459	-9.428	1.00	0.21
MOTA	1536	HB	THR	105	15.018	6.233	-8.704	1.00	0.21
ATOM	1537	OG1	THR	105	15.806		-10.447	1.00	0.24
ATOM	1538	HG1	THR	105	15.882	6 404	-10.752	1.00	
ATOM	1539	CG2			14.846	4.101	-8.729		0.86
ATOM		HG21		105	15.178			1.00	0.21
ATOM		HG22		105		4.233	-7.711	1.00	1.04
ATOM	1542	HG23	THR		15.524	3.442	-9.249	1.00	1.07
ATOM	1543			105	13.854	3.674	-8.731	1.00	0.99
MOTA		C	THR	105	13.166	4.597	-11.087	1.00	0.23
	1544	0	THR	105	12.521	3.606	-10.808	1.00	0.23
MOTA	1545	N	SER	106	13.668	4.769	-12.282	1.00	0.26
ATOM	1546	HN	SER	106	14.194	5.572	-12.480	1.00	0.29
MOTA	1547	CA	SER	106	13.454		-13.337	1.00	0.29
MOTA	1548	HA	SER	106	12.570	3.163	-13.111	1.00	0.30
ATOM	1549	CB	SER	106	13.290		-14.695	1.00	0.35
MOTA	1550	HB1	SER	106	14.249	4.467		1.00	
MOTA	1551	HB2	SER	106	12.916				1.09
MOTA	1552	OG	SER	106	12.365		-14.554	1.00	0.96
ATOM	1553	HG	SER	106			-15.483	1.00	1.44
ATOM	1554	C	SER		11.671	4.285	-15.766	1.00	1.97
ATOM	1555			106	14.674	2.817	-13.372	1.00	0.28
ATOM		0	SER	106	14.669	1.781	-14.006	1.00	0.31
	1556	N	SER	107	15.715	3.187	-12.677	1.00	0.26
MOTA	1557	HN	SER	107	15.687	4.023	-12.166	1.00	0.25
ATOM	1558	CA	SER	107	16.940	2.340	-12.641	1.00	0.27
ATOM	1559	HA	SER	107	17.018		-13.560	1.00	0.29
ATOM	1560	CB	SER	107	18.175	3.226	-12.474	1.00	0.28
MOTA	1561	HB1	SER	107	18.292	3 847	-13.353	1.00	1.12
MOTA	1562	HB2	SER	107	19.049		-12.355		
MOTA	1563	OG	SER	107	18.017	4 040	-11.320	1.00	1.04
MOTA	1564	HG	SER	107	18.556	4 007	-11.320	1.00	1.29
MOTA	1565	c	SER			4.827	-11.436	1.00	1.82
ATOM	1566	ŏ	SER	107	16.836	1.376	-11.460	1.00	0.26
ATOM	1567			107	15.829	1.324	-10.781	1.00	0.26
		N	SER	108	17.859	0.609	-11.203	1.00	0.28
MOTA	1568	HN	SER	108	18.666	0.658	-11.757	1.00	0.31
MOTA	1569	CA	SER	108	17.788	-0.342	-10.061	1.00	0.30
ATOM	1570	HA	SER	108	16.775	-0.706	-9.967	1.00	0.30
MOTA	1571	CB	SER	108	18.728	-1.527	-10.330	1.00	0.36
MOTA	1572	HB1	SER	108	19.561	-1.505	-9.642		
MOTA	1573	HB2	SER	108	19.103		-11.338	1.00	1.09
ATOM	1574	OG	SER	108	18.005			1.00	0.95
MOTA	1575	HG	SER	108	18.550		-10.176	1.00	1.47
MOTA	1576	c	SER	108			-10.513	1.00	2.00
ATOM	1577	ŏ	SER	108	18.181	0.390	-8.767	1.00	0.28
ATOM	1578	N	LYS		19.279	0.265	-8.261	1.00	0.33
ATOM	1579			109	17.272	1.157	-8.224	1.00	0.24
	-3/3	HN	LYS	109	16 302	1 241	0 646	1 00	^ ^^

ATOM	1580	CA	LYS	109	17.561	1.897	-6.960	1.00	0.23
ATOM	1581	HA	LYS	109	18.275	1.341	-6.370	1.00	0.25
ATOM	1582		LYS	109	18.123		_		
		HB1				3.293	-7.268	1.00	0.24
MOTA	1583			109	18.172	3.868	-6.355	1.00	0.27
MOTA	1584	HB2		109	17.472	3.793	-7.970	1.00	0.25
ATOM	1585	CG	LYS	109	19.525	3.177	-7.868	1.00	0.30
MOTA	1586	HG1	LYS	109	19.476	2.615	-8.785	1.00	0.54
ATOM	1587	HG2		109	20.177	2.675	-7.170	1.00	0.70
ATOM	1588		LYS	109					
					20.072	4.574	-8.169	1.00	0.75
MOTA	1589	HD1		109	20.124	5.144	-7.254	1.00	1.27
MOTA	1590	HD2		109	19.420	5.074	-8.870	1.00	1.27
ATOM:	1591	CE	LYS	109	21.475	4.453	-8.770	1.00	1.13
MOTA	1592	HE1	LYS	109	21.396	4.264	-9.830	1.00	1.68
MOTA	1593	HE2	LYS	109	22.000	3.636	-8.297	1.00	1.68
MOTA	1594		LYS	109	22.224	5.721	-8.545		
ATOM	1595	HZ1		109				1.00	1.79
MOTA	1596				21.689	6.516	-8.948	1.00	2.22
		HZ2		109	23.155	5.660	-9.006	1.00	2.17
MOTA	1597	HZ3		109	22.351	5.873	-7.525	1.00	2.34
MOTA	1598	C	LYS	109	16.259	2.052	-6.175	1.00	0.21
ATOM	1599	0	LYS	109	15.190	2.110	-6.747	1.00	0.20
ATOM	1600	N	GLY	110	16.338	2.124	-4.873	1.00	0.23
ATOM	1601		GLY	110	17.212	2.079	-4.432		
ATOM	1602		GLY	110				1.00	0.26
					15.099	2.283	-4.056	1.00	0.22
ATOM	1603		GLY	110	14.751	3.302	-4.124	1.00	0.23
MOTA	1604		GLY	110	15.316	2.044	-3.024	1.00	0.25
MOTA	1605	C	GLY	110	14.013	1.342	-4.581	1.00	0.19
ATOM	1606	0	GLY	110	14.281	· 0.216	-4.949	1.00	0.20
ATOM	1607	N	TYR	111	12.789	1.801	-4.626	1.00	0.17
MOTA	1608		TYR	111	12.599	2.716			
ATOM	1609	CA	TYR	111			-4.330	1.00	0.18
					11.683	0.941	-5.136	1.00	0.15
MOTA	1610		TYR	111	11.975	-0.098	-5.088	1.00	0.16
MOTA	1611	CB	TYR	111	10.437	1.162	-4.277	1.00	0.15
MOTA	1612	HB1	TYR	111	9633	0.540	-4.641	1.00	0.15
MOTA	1613	HB2	TYR	111	10.143	2.200	-4.330	1.00	0.16
ATOM	1614	CG	TYR	111	10.745	0.798	-2.844	1.00	0.17
MOTA	1615	CD1		111	10.648	-0.533	-2.422	1.00	
ATOM	1616		TYR	111				_	0.17
					10.354	-1.301	-3.121	1.00	0.17
ATOM	1617	CD2	TYR	111.	11.127	1.794	-1.936	1.00	0.20
ATOM	1618	HD2	TYR	111	11.201	2.821	-2.261	1.00	0.23
MOTA	1619	CE1	TYR	111	10.933	-0.868	-1.093	1.00	0.19
MOTA	1620	HE1	TYR	111	10.858	-1.895	-0.767	1.00	0.20
ATOM	1621	CE2	TYR	111	11.412	1.459	-0.607	1.00	0.22
ATOM	1622	HE2	TYR	111	11.706	2.227			
ATOM	1623	CZ	TYR	111			0.093	1.00	0.26
ATOM					11.315	0.127	-0.185	1.00	0.21
	1624	OH	TYR	111	11.595	-0.204	1.125	1.00	0.23
ATOM	1625	HH	TYR	111	12.543	-0.121	1.255	1.00	0.95
ATOM	1626	C	TYR	111	11.374	1.321	-6.588	1.00	0.14
ATOM	1627	0	TYR	111	10.949	2.424	-6.871	1.00	0.15
ATOM	1628	N	ASN	112	11.581	0.421	-7.511	1.00	0.15
MOTA	1629	HN	ASN	112	11.924	-0.464	-7.264	1.00	0.17
MOTA	1630	CA	ASN	112	11.295	0.739		1.00	
ATOM	1631	HA	ASN				-8.939	1.00	0.16
				112		1.605	-9.235	1.00	0.16
MOTA	1632	CB	ASN	112	11.677	-0.450	-9.822	1.00	0.19
MOTA	1633	HB1		112	11.025	-1.276	-9.607	1.00	0.22
MOTA	1634	HB2		112	12.698	-0.739	-9.622	1.00	0.19
MOTA	1635	CG	asn	112	11.531	-0.060	-11.295	1.00	0.24
MOTA	1636	OD1	asn	112	10.446	0.248	-11.748	1.00	0.96
MOTA	1637	ND2	ASN	112	12.583		-12.067	1.00	1.06
MOTA	1638	HD21	ASN	112	13.458		-11.704	1.00	1.80
MOTA	1639	HD22	ASN	112	12.497	0.300	-13.012		
MOTA	1640	c	ASN	112				1.00	1.08
ATOM					9.803	1.040	-9.108	1.00	0.15
	1641	0	ASN	112	8.953	0.310	-8.637	1.00	0.14
MOTA	1642	N	LEU	113	9.482	2.112	-9.777	1.00	0.15
MOTA	1643	HN	LEU	113	10.187	2.684	-10.145	1.00	0.16
ATOM	1644	CA	LEU	113	8.049	2.475	-9.984	1.00	0.15
ATOM	1645	HA	LEU	113	7.582	2.620	-9.025	1.00	0.14
MOTA	1646	CB	LEU	113	7.981				
MOTA	1647	HB1		113			-10.791	1.00	0.16
					8.513		-11.721	1.00	0.17
MOTA	1648	HB2		113	8.452		-10.226	1.00	0.16
ATOM	1649	CG	LEU	113	6.523		-11.095	1.00	0.17
ATOM	1650	HG	LEU	113	6.041		-11.652	1.00	0.18
MOTA	1651	CD1		113	5.748	4.421	-9.793	1.00	0.18
MOTA	1652	HD11	LEU	113	4.841		-10.007	1.00	0.99
ATOM		HD12		113	6.359	4.991			
ATOM		HD13		113	5.490		-9.110	1.00	1.00
ATOM	1655	CD2		113		3.474	-9.343	1.00	0.97
ATOM			I.EII	113	6.526	5.457	-11.943	1.00	0.20
n. um	7030	DUZ I	1 . P.1 1	114	6 115	E 777	-11 274	1 00	1 05

ATOM	1657	HD22	LEU	113	5.930	5 302	-12.830	1 00	1 02
MOTA	1658			113	7.539		-12.231	1.00	1.03
MOTA	1659	С	LEU	113	7.320	1 361	-10.743		1.00
MOTA	1660	ō	LEU	113	6.203			1.00	0.15
ATOM	1661	Ŋ	PHE	114	7.928		-10.419	1.00	0.15
ATOM	1662	HN	PHE	114			-11.762	1.00	0.16
ATOM	1663	CA	PHE		8.822	1.123	-12.020	1.00	0.17
ATOM				114	7.245	-0.250	-12.555	1.00	0.17
	1664	HA	PHE	114	6.338	0.151	-12.980	1.00	0.18
ATOM	1665	CB	PHE	114	8.159	-0.720	-13.685	1.00	0.21
ATOM	1666	HB1		114	9.077	-1.108	-13.271	1.00	0.22
ATOM	1667	HB2		114	8.380	0.111	-14.340	1.00	0.22
MOTA	1668	CG	PHE	114	7.457	-1.807	-14.464	1.00	0.24
MOTA	1669	CD1	PHE	114	7.545	-3.135		1.00	0.35
MOTA	1670	HD1	PHE	114	8.105		-13.147	1.00	0.43
MOTA	1671	CD2	PHE	114	6.724	-1.494	-15.613	1.00	0.24
ATOM	1672	HD2		114	6.655	-0 470	-15.950		
MOTA	1673	CE1		114	6.902	-4.140	-14.741	1.00	0.28
MOTA	1674	HE1		114	6.975	-5.171		1.00	0.39
ATOM	1675	CE2		114			-14.402	1.00	0.50
ATOM	1676	HE2		114	6.078	-2.512	-16.327	1.00	0.26
ATOM	1677				5.511	-2.273	-17.214	1.00	0.30
		CZ	PHE	114	6.168	-3.839	-15.890	1.00	0.32
MOTA	1678	HZ	PHE	114	5.670	-4.623	-16.438	1.00	0.35
ATOM	1679	C	PHE	114	6.900	-1.452	-11.676	1.00	0.17
ATOM	1680	0	PHE	114	5.842	-2.034	-11.806	1.00	0.17
MOTA	1681	N	LEU	115	7.774	-1.846	-10.797	1.00	0.18
MOTA	1682	HN	LEU	115	8.631		-10.706	1.00	0.18
atom	1683	CA	LEU	115	7.463	-3.028	-9.946	1.00	0.20
ATOM	1684	HA	LEU	115	7.297		-10.579	1.00	0.21
MOTA	1685	CB	LEU	115	8.634	-3.304	-8.984		
ATOM	1686		LEU	115	8.237	-3.650		1.00	0.23
MOTA	1687	HB2		115	9.172		-8.041	1.00	0.26
MOTA	1688	CG	LEU	115	9.612	-2.387	-8.821	1.00	0.22
ATOM	1689	HG	LEU	115		-4.369	-9.539	1.00	0.28
ATOM	1690				10.397	-4.525	-8.812	1.00	0.33
		17011	LEU	115	8.886	-5.702	-9.749	1.00	0.36
MOTA		HD11		115	9.551	-6.514	-9.498	1.00	0.99
ATOM	1692	HD12		115	8.578	-5.795	-10.779	1.00	1.11
ATOM	1693			115	8.017	-5.740	-9.109	1.00	1.13
MOTA	1694		LEU	115	10.249	-3.903	-10.859	1.00	0.30
ATOM	1695	HD21	LEU	115	10.497	-4.761		1.00	1.10
ATOM	1696	HD22	LEU	115	11.149		-10.645	1.00	1.06
ATOM	1697	HD23	LEU	115	9.567	-3.272	-11.395	1.00	
MOTA	1698	C	LEU	115	6.194	-2.748			1.01
ATOM	1699	ō	LEU	115	5.280		-9.136	1.00	0.19
ATOM	1700	N	VAL	116	6.130	-3.548	-9.106	1.00	0.20
ATOM	1701	HN	VAL	116		-1.624	-8.475	1.00	0.18
ATOM	1702	CA	VAL	116	6.879	-0.993	-8.508	1.00	0.18
ATOM	1703	HA			4.919	-1.305	-7.664	1.00	0.19
ATOM	1704		VAL	116	4.686	-2.146	-7.028	1.00	0.21
		CB	VAL	116	5.203	-0.078	-6.794	1.00	0.20
MOTA	1705	HB	VAL	116	5.581	0.722	-7.414	1.00	0.19
ATOM	1706	CG1	VAL	116	3.914	0.381	-6.103	1.00	0.22
MOTA		HG11		116	3.253	0.832	-6.828	1.00	1.05
MOTA		HG12		116	4.155	1.105	-5.339	1.00	1.05
MOTA	1709	HG13	VAL	116	3.426	-0.470	-5.650	1.00	1.03
ATOM	1710	CG2	VAL	116	6.246	-0.443	-5.737	1.00	0.21
MOTA	1711	HG21	VAL	116	7.188	-0.654	-6.221		
MOTA	1712	HG22	VAL	116	5.917	-1.317	-5.194	1.00	1.02
ATOM	1713	HG23	VAL	116	6.370			1.00	0.98
MOTA	1714	C	VAL	116	3.724	0.382	-5.052	1.00	1.03
MOTA	1715	ō	VAL	116		-1.020	-8.582	1.00	0.18
ATOM	1716	N	ALA	117	2.615	-1.433	-8.312	1.00	0.19
ATOM	1717	HN			3.934	-0.307	-9.659	1.00	0.17
ATOM	1718		ALA	117	4.833	0.028	-9.859	1.00	0.16
ATOM		CA	ALA	117	2.796	0.007	-10.572	1.00	0.17
	1719	HA	ALA	117	2.064	0.598	-10.044	1.00	0.19
MOTA	1720	CB	ALA	117	3.306	0.795	-11.780	1.00	0.18
MOTA	1721		ALA	117	4.378	0.709	-11.840	1.00	1.05
MOTA	1722	HB2	ALA	117	3.033		-11.674	1.00	1.01
MOTA	1723	HB3	ALA	117	2.863		-12.682	1.00	0.98
ATOM	1724	С	ALA	117	2.150		-11.058		
MOTA	1725	ŏ	ALA	117	0.956			1.00	0.17
MOTA	1726	N	ALA	118		-1.450	-10.951	1.00	0.19
MOTA	1727	HN	ALA	118	2.931	-2.187	-11.588	1.00	0.16
MOTA	1728	CA	ALA		3.893		-11.663	1.00	0.16
ATOM	1729		ALA	118	2.366		-12.083	1.00	0.17
ATOM		HA		118	1.643	-3.273	-12.859	1.00	0.19
	1730	CB	ALA	118	3.491	-4.335	-12.653	1.00	0.17
ATOM	1731		ALA	118	3.125	-5.338	-12.812	1.00	1.05
MOTA	1732	_	ALA	118	4.316	-4.358	-11.956	1.00	1.02
MOTA	1733	HB3	ALA	118	3 824	-3 03N	_13 503	1 00	1 72

MOTA	1734	С	ALA	118	1.687	-4 220	-10.935	1 00	0 17
ATOM	1735	ŏ	ALA	118	0.699		-11.124	1.00	0.17 0.18
ATOM	1736	N	HIS	119	2.225	-4.123	-9.751	1.00	0.16
ATOM	1737	HN	HIS	119	3.035	-3.585	-9.623	1.00	0.16
MOTA	1738	CA	HIS	119	1.627	-4.855	-8.599	1.00	0.17
MOTA	1739	HA	HIS	119	1.576	-5.907	-8.833	1.00	0.18
ATOM	1740	CB	HIS	119	2.513	-4.655	-7.368	1.00	0.19
MOTA	1741	HB1	HIS	119	2.547	-3.605	-7.116	1.00	0.19
MOTA	1742	HB2	HIS	119	3.512	-5.005	-7.584	1.00	0.20
MOTA	1743	CG	HIS	119	1.950	-5.431	-6.210	1.00	0.21
MOTA	1744		HIS	119	2.228	-6.775	-6.020	1.00	0.26
MOTA	1745		HIS	119	2.791	-7.336	-6.593	1.00	0.30
MOTA	1746		HIS	119	1.128	-5.067	-5.172	1.00	0.20
MOTA	1747		HIS	119	0.719	-4.079	-5.019	1.00	0.21
MOTA	1748		HIS	119	1.585	-7.168	-4.906	1.00	0.27
MOTA ATOM	1749		HIS	119	1.622	-8.171	-4.509	1.00	0.33
MOTA	1750 1751		HIS	119	0.899	-6.166	-4.350	1.00	0.23
ATOM	1752	CO	HIS HIS	119 119	0.215 -0.721	-4.333	-8.299	1.00	0.17
ATOM	1753	N	GLU	120		-5.101	-8.185	1.00	0.18
ATOM	1754	HN	GLU	120	0.043 0.801	-3.044 $-2.430$	-8.160	1.00	0.18
ATOM	1755	CA	GLU	120	-1.322	-2.520	-8.248 -7.860	1.00	0.18
ATOM	1756	HA	GLU	120	-1.666	-2.977	-6.943	1.00	0.20 0.21
ATOM	1757	СВ	GLU	120	-1.294	-0.999	-7.668	1.00	0.21
MOTA	1758		GLU	120	-0.719	-0.763	-6.785	1.00	0.37
ATOM	1759			120	-2.302	-0.635	-7.542	1.00	0.33
MOTA	1760	CG	GLU	120	-0.663	-0.314	-8.875	1.00	0.41
MOTA	1761	HG1	GLU	120	-1.125	-0.668	-9.781	1.00	0.63
MOTA	1762	HG2	GLU	120	0.393	-0.531	-8.895	1.00	0.87
MOTA	1763	ÇD	GLU	120	-0.875	1.194	-8.757	1.00	0.94
MOTA	1764	OE1	GLU	120	-0.757	1.703	-7.654	1.00	1.67
MOTA	1765	OE2		120	-1.151	1.816	-9.769	1.00	1.56
ATOM	1766	C	GLU	120	-2.291	-2.903	-8.984	1.00	0.20
MOTA	1767	0	GLU	120	-3.432	-3.238	-8.737	1.00	0.21
ATOM	1768	N	PHE	121	-1.853	-2.872	-10.217	1.00	0.19
ATOM	1769	HN	PHE	121	-0.928		-10.405	1.00	0.19
ATOM	1770	CA	PHE	121	-2.767		-11.331	1.00	0.21
MOTA	1771	HA	PHE	121	-3.628		-11.317	1.00	0.23
MOTA	1772	CB	PHE	121	-2.053		-12.685	1.00	0.22
ATOM ATOM	1773 1774	HB1		121	-2.576		-13.419	1.00	0.24
ATOM	1775	HB2 CG		121 121	-1.041		-12.587	1.00	0.21
ATOM	1776		PHE	121	-2.026		-13.141	1.00	0.25
ATOM	1777		PHE	121	-0.804 0.121		-13.308 -13.113	1.00	0.27
ATOM	1778		PHE	121	-3.227		-13.113	1.00	0.40
MOTA	1779		PHE	121	-4.173		-13.281	1.00	0.45 0.60
ATOM	1780		PHE	121	-0.781		-13.733	1.00	0.29
ATOM	1781		PHE	121	0.163		-13.862	1.00	0.39
MOTA	1782	CE2	PHE	121	-3.202		-13.828	1.00	0.49
MOTA	1783	HE2		121	-4.127		-14.029	1.00	0.68
MOTA	1784	CZ	PHE	121	-1.979	0.988	-13.993	1.00	0.34
ATOM	1785	HZ	PHE	121	-1.961		-14.321	1.00	0.38
MOTA	1786	C	PHE	121	-3.228		-11.120	1.00	0.20
MOTA MOTA	1787	0	PHE	121	-4.374		-11.344	1.00	0.21
MOTA	1788 1789	N	GLY	122	-2.344		-10.690	1.00	0.18
MOTA	1790	HN CA	GLY GLY	122 122	-1.424		-10.514	1.00	0.17
MOTA	1791	HAI		122	-2.737 -1.890	7 522	-10.464	1.00	0.20
ATOM	1792	HA2		122	-3.072	-7.523	-10.092 -11.394	1.00	0.21
MOTA	1793	c	GLY	122	-3.867	-7.022	-9.435	1.00	0.21
ATOM	1794	ō	GLY	122	-4.823	<b>-7.756</b>	-9.589	1.00	0.20 0.22
ATOM	1795	Ň	HIS	123	-3.778	-6.240	-8.392	1.00	0.20
ATOM	1796	HN	HIS	123	-3.005	-5.644	-8.287	1.00	0.20
MOTA	1797	CA	HIS	123	-4.864	-6.243	-7.371	1.00	0.22
MOTA	1798	HA	HIS	123	-5.047	-7.255	-7.042	1.00	0.23
ATOM	1799	CB	HIS	123	-4.456	-5.382	-6.174	1.00	0.25
ATOM	1800		HIS	123	-5.324	-5.180	-5.564	1.00	0.30
MOTA	1801		HIS	123	-4.041	-4.449	-6.527	1.00	0.25
MOTA	1802	CG	HIS	123	-3.427	-6.108	-5.354	1.00	0.27
MOTA	1803		HIS	123	-3.736	-7.247	-4.628	1.00	0.37
MOTA	1804		HIS	123	-4.611	-7.685	-4.581	1.00	0.45
MOTA MOTA	1805 1806		HIS	123	-2.096	-5.866	-5.125	1.00	0.25
ATOM	1806	בעת רפי	HIS	123	-1.532	-5.046		1.00	0.27
ATOM	1807		HIS	123 123	-2.614	-7.644	-4.001	1.00	0.38
MOTA	1809		HIS	123	-2.553 -1.584	-8.514 -6.837	-3.367	1.00	0.47
MOTA	1810	C	HIS	123	-F 137	-0.83/ -5 671	-4.269 -7 003	1.00	0.29
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MOTA	1811	0	HIS	123	-7.229	-6.148	-7.755	1.00	0.25
MOTA	1812	N	SER	124	-6.002	-4.646	-8.788	1.00	0.23
MOTA	1813	HN	SER	124	-5.110	-4.278	-8.962	1.00	0.22
MOTA	1814	CA	SER	124	-7.196	-4.030	-9.429	1.00	0.25
ATOM	1815	HA	SER	124	-7.928	-3.790	-8.672	1.00	0.27
MOTA	1816	CB	SER	124	-6.778		-10.156	1.00	0.27
MOTA	1817	HB1	SER	124	-6.219	-2.119	-9.478	1.00	0.29
ATOM	1818	HB2	SER	124	-7.654	-2.224	-10.494	1.00	0.29
MOTA	1819	OG	SER	124	-5.975		-11.279	1.00	0.25
MOTA	1820	HG	SER	124	-6.545		-12.050	1.00	0.88
ATOM	1821	C.	SER	124	-7.805	-5.006	-10.437	1.00	0.24
ATOM	1822	0	SER	124	-8.975	-4.932	-10.755	1.00	0.26
ATOM	1823	N	LEU	125			-10.952		
					-7.022			1.00	0.22
ATOM	1824	HN	LEU	125	-6.078		-10.690	1.00	0.21
MOTA	1825	CA	LEU	125	-7.562	-6.879	-11.949	1.00	0.23
MOTA	1826	HA	LEU	125	-8.285		-12.568	1.00	0.24
MOTA	1827	CB	LEU	125	-6.420		-12.827	1.00	0.22
MOTA	1828	HB1	LEU	125	-6.759	-8.247	-13.398	1.00	0.24
MOTA	1829	HB2	LEU	125	-5.594	-7.698	-12.197	1.00	0.22
ATOM	1830	CG	LEU	125			-13.779		
					-5.956			1.00	0.22
ATOM	1831	HG	LEU	125	-5.928		-13.241	1.00	0.24
MOTA	1832	CD1	LEU	125	-4.556	-6.601	-14.302	1.00	0.25
MOTA		HD11		125	-4.588		-14.874	1.00	0.99
MOTA		HD12		125	-3.879		-13.471	1.00	1.00
MOTA	1835	HD13	LEU	125	-4.215	-5.794	-14.933	1.00	1.05
ATOM	1836	CD2	LEU	125	-6.913		-14.976	1.00	0.24
ATOM		HD21	LEU	125	-7.793		-14.682		
								1.00	1.05
MOTA		HD22	LEU	125	-7.201	-7.135		1.00	1.00
MOTA	1839	HD23	LEU	125	-6.415	-5.627	-15.775	1.00	1.03
MOTA	1840	С	LEU	125	-8.256		-11.234	1.00	0.24
MOTA	1841	0	LEU	125	-8.790		-11.864	1.00	0.33
MOTA	1842	N	GLY	126	-8.277	-8.035	-9.927	1.00	0.24
ATOM	1843	HN	GLY	126	-7.858	-7.298	-9.435	1.00	0.29
ATOM	1844	CA	GLY	126					
					-8.968	-9.132	-9.185	1.00	0.27
MOTA	1845	HAL	GLY	126	-9.748	-9.545	-9.807	1.00	0.29
ATOM	1846	HA2	GLY	126	-9.408	-8.727	-8.285	1.00	0.29
ATOM	1847	С	GLY	126		-10.245	-8.809		
								1.00	0.26
ATOM	1848	0	GLY	126		-11,268	-8.283	1.00	0.30
MOTA	1849	N	LEU	127	-6.719	-10.068	-9.063	1.00	0.23
MOTA	1850	HN	LEU	127	-6.410	-9.239	-9.484	1.00	0.22
MOTA	1851	CA	LEU	127		-11.138	-8.700	1.00	0.25
MOTA	1852	HA	LEU	127	-6.212	-12.099	-8.815	1.00	0.28
ATOM	1853	CB	LEU	127	-4.507	-11.052	-9.602	1.00	0.23
MOTA	1854		LEU	127					
						-11.696		1.00	0.25
MOTA	1855	HB2	LEU	127	-4.156	-10.033	-9.602	1.00	0.22
ATOM	1856	CG	LEU	127	-4.844	-11.471	-11.045	1.00	0.24
ATOM	1857	HG	LEU	127			-11.384	1.00	0.23
ATOM	1858		LEU	127			-11.962	1.00	0.24
MOTA	1859	HD11	LEU	127	-4.001	-10.692	-12.868	1.00	1.00
ATOM	1860	HD12	LEU	127		-12.073		1.00	1.02
MOTA		HD13		. 127			-11.460		
								1.00	1.03
MOTA	1862		LEU	127	-5.150	-12.980	-11.109	1.00	0.30
MOTA	1863	HD21	LEU	127	-5.021	-13.334	-12.121	1.00	1.04
ATOM	1864	HD22	LEU	127	-6 169	-13.150	-10.805	1.00	1.11
ATOM			LEU	127					
						-13,515		1.00	1.03
MOTA	1866	C	LEU	127	-5.315	-10.969	-7.241	1.00	0.28
MOTA	1867	0	LEU	127	-5.245	-9.872	-6.723	1.00	0.32
MOTA	1868	N	ASP	128	-5 027	-12.059	-6.581	1.00	0.32
MOTA	1869	HN							
			ASP	128		-12.928	-7.029	1.00	0.34
MOTA	1870	ÇA	ASP	128	-4.598	-11.997	-5.154	1.00	0.39
ATOM	1871	HA	ASP	128	-4.882	-11.046	-4.728	1.00	0.40
MOTA	1872	CB	ASP	128		-13.130			
								1.00	0.48
ATOM	1873		ASP	128		-14.064		1.00	0.48
ATOM	1874	HB2	ASP	128	-6.311	-13.193	-4.661	1.00	0.50
MOTA	1875	CG	ASP	128	-5 171	-12.854	-2.873		0.55
								1.00	
MOTA	1876		ASP	128		-12.980		1.00	1.23
ATOM	1877	QD2	ASP	128	-6.185	-12.521	-2.283	1.00	1.22
MOTA	1878	С	ASP	128		-12.159		1.00	0.37
ATOM									
	1879	0	ASP	128		-12.387		1.00	0.59
MOTA	1880	N	HIS	129	-2.507	-12.042	-3.914	1.00	0.23
MOTA	1881	HN	HIS	129		-11.856		1.00	0.32
ATOM	1882	CA	HIS	129	_1 020	-12.189	2.110		
								1.00	0.22
ATOM	1883	HA	HIS	129		-11.439		1.00	0.21
ATOM	1884	CB	HIS	129	-0.606	-12.019		1.00	0.23
ATOM	1885		HIS	129		-12.302		1.00	0.24
ATOM									
	1886		HIS	129		-12.653		1.00	0.25
MOTA	1887	CG	HIS	129	-0.779	-10.585	-1.912	1.00	0.22
							_		

MOTA	1888	ND1 HIS	129	-1.862	-10.161	-1.156	1.00	0.35
MOTA	1889	HD1 HIS	129	-2,602	-10.720	-0.841	1.00	0.53
MOTA	1890	CD2 HIS	129	-0.007	-9.468	-2.118	1.00	0.34
MOTA	1891	HD2 HIS	129	0.918	-9.447	-2.673	1.00	
ATOM	1892	CE1 HIS	129	-1.711				0.54
	1893	HE1 HIS			-8.842	-0.936	1.00	0.31
ATOM			129	-2.406	-8.239	-0.370	1.00	0.44
MOTA	1894	NE2 HIS	129	-0.597	-8.369	-1.501	1.00	0.28
MOTA	1895	C HIS	129		-13.584	-4.277	1.00	0.24
MOTA	1896	O HIS	129	-1.267	-14.568	-3.991	1.00	0.28
ATOM	1897	N SER	130	0.474	-13.671	-4.999	1.00	0.24
MOTA	1898	HN SER	130	0.984	-12.862	-5.210	1.00	0.23
ATOM	1899	CA SER	130		-14.996	-5.498	1.00	0.29
ATOM	1900	HA SER	130		-15.710	-5.464	1.00	0.33
MOTA	1901	CB SER	130		-14.852		1.00	
ATOM			130			-6.938		0.32
	1902				-14.082	-6.982	1.00	0.31
MOTA	1903	HB2 SER	130		-14.577	-7.576	1.00	0.35
MOTA	1904	OG SER	130		-16.092	-7.378	1.00	0.40
ATOM	1905	HG SER	130		-16.714	-7.469	1.00	0.97
MOTA	1906	C SER	130		-15.484	-4.609	1.00	0.28
MOTA	1907	O SER	130	2.801	-14.696	-4.009	1.00	0.29
MOTA	1908	N LYS	131	2.287	-16.775	-4.514	1.00	0.30
MOTA	1909	HN LYS	131	1.705	-17.393	-5.003	1.00	0.32
MOTA	1910	CA LYS	131	3.386	-17.310	-3.656	1.00	0.32
MOTA	1911	HA LYS	131	3.665	-16.567	-2.923	1.00	0.34
ATOM	1912	CB LYS	131	2 903	-18.572	-2.936	1.00	0.39
ATOM	1913	HB1 LYS	131	3 714	-18.988	-2.355		
ATOM	1914	HB2 LYS	131	2.714	-19.298,		1.00	0.42
	1915			2.5/2	-19.298	-3.664	1.00	0.40
ATOM		CG LYS	131	. 1.743	-18.214	-2.003	1.00	0.45
ATOM	1916	HG1 LYS	131		-17.798	-2.581	1.00	0.79
ATOM	1917	HG2 LYS	131	2.077	-17.488	-1.276	1.00	1.01
MOTA	1918	CD LYS	131		-19.472	-1.280	1.00	1.18
MOTA	1919	HD1 LYS	131	2.064	-19.890	-0.698	1.00	1.86
MOTA	1920	HD2 LYS	131	0.921	-20.199	-2.006	1.00	1.66
MOTA	1921	CE LYS	131	0.096	-19.108	-0.349	1.00	1.52
MOTA	1922	HE1 LYS	131	-0.788	-18.908	-0.937	1.00	1.92
MOTA	1923	HE2 LYS	131	0.355	-18.229	0.222	1.00	1.93
ATOM	1924	NZ LYS	131	-0.174	-20.242	0.581	1.00	2.23
ATOM	1925	HZ1 LYS	131	-1.103	-20.109	1.030	1.00	2.72
MOTA	1926	HZ2 LYS	131	0.565	-20.272	1.313		
ATOM	1927	HZ3 LYS	131		-21.135		1.00	2.53
MOTA	1928	C LYS	131	-0.174	-17.649	0.050	1.00	2.72
MOTA	1929		131	4.604	-17.649	-4.521	1.00	0.31
				5.612	-18.116	-4.027	1.00	0.34
MOTA	1930	N ASP	132		-17.411	-5.804	1.00	0.29
MOTA	1931	HN ASP	132	3.717	-17.028	-6.190	1.00	0.28
MOTA	1932	CA ASP	132 ·		-17.719	-6.674	1.00	0.30
MOTA	1933	HA ASP	132	6.187		-6.302	1.00	0.32
MOTA	1934	CB ASP	132	5.225	-17.970	-8.108	1.00	0.32
ATOM	1935	HB1 ASP	132	4.727	-17.090	-8.483	1.00	0.31
MOTA	1936	HB2 ASP	132	4.539	-18.804	-8.118	1.00	0.34
MOTA	1937	CG ASP	132		-18.289	-8.996	1.00	0.35
MOTA	1938	OD1 ASP	132		-19.371	-9.558	1.00	1.10
MOTA	1939	OD2 ASP	132		-17.446		1.00	1.15
ATOM	1940	C ASP	132	6.656	-16.501	-6.659	1.00	0.28
ATOM	1941	O ASP	132		-15.399	-6.939	1.00	0.28
MOTA	1942	N PRO	133	7.930	-16.658	-6.328	1.00	0.30
MOTA	1943	CA PRO	133		-15.484	-6.296	1.00	0.31
MOTA	1944	HA PRO	133		-14.766	-5.566		0.32
ATOM	1945	CB PRO	133	10 173	-16.097		1.00	
ATOM	1946	HB1 PRO	133		-15.694	-5.832	1.00	0.36
ATOM	1947	HB2 PRO	133	10.441	-15.094	-4.867	1.00	0.36
ATOM	1948				-15.869	-6.549	1.00	0.41
		CG PRO	133		-17.615	-5.721	1.00	0.42
MOTA	1949	HG1 PRO	133		-17.940	-4.732	1.00	0.51
MOTA	1950	HG2 PRO	133		-18.103	-6.457	1.00	0.51
MOTA	1951	CD PRO	133		-17.972	-5.969	1.00	0.35
MOTA	1952	HD2 PRO	133	8.456	-18.679	-6.785	1.00	0.34
MOTA	1953	HD1 PRO	133	8.091	-18.362	-5.069	1.00	0.38
MOTA	1954	C PRO	133		-14.810	-7.662	1.00	0.31
MOTA	1955	O PRO	133		-13.691	-7.749	1.00	0.34
MOTA	1956	N GLY	134		-15.477	-8.729	1.00	0.32
MOTA	1957	HN GLY	134		-16.382	-8.647	1.00	0.35
ATOM	1958	CA GLY	134	8 860	-14.856	-10 074	1.00	0.34
ATOM	1959	HA1 GLY	134	2.000	-15.630	-10.074	1.00	0.37
ATOM	1960	HA2 GLY	134	9.040	-14.177	-10.003		
MOTA	1961	C GLY	134	7 E00	-14.087	-10.047	1.00	0.36
MOTA	1962	O GLY	134	7.578	-12 400	-10.4/1	1.00	0.29
MOTA	1963	N ALA	135	1.203	-13.420		1.00	0.29
ATOM	1964	HN ALA			-14.168	-9.683	1.00	0.27
AI OF	2504	· ··· ·	135	6.607	-14.709	-8.867	1 00	N.28

MOTA	1965	CA	ALA	135	5 312	-13.434	-10 026	1 00	0.24
ATOM	1966	HA	ALA	135	5.312			1.00	
MOTA	1967		_				-11.099	1.00	0.25
		CB	ALA	135		-14.151	-9.410	1.00	0.25
MOTA	1968	HB1		135	3.633	-14.765	-10.160	1.00	1.07
MOTA	1969	HB2	ALA	135	3.405	-13.421	-9.041	1.00	1.01
ATOM	1970	HB3	ALA	135	4.442	-14.774	-8.593	1.00	1.04
ATOM	1971	C	ALA	135	5 388	-12.007	-9.479	1.00	0.21
ATOM	1972	ŏ	ALA	135	5.500	11.7007			
					3.908	-11.760	-8.440	1.00	0.23
MOTA	1973	N	LEU	136	4.799	-11.067	-10.164	1.00	0.22
MOTA	1974	HN	LEU	136	4.330	-11.286	-10.996	1.00	0.24
MOTA	1975	CA	LEU	136	4.830	-9.660	-9.676	1.00	0.23
MOTA	1976	HA	LEU	136	5.842	-9.382	-9.427	1.00	0.25
MOTA	1977	СВ	LEU	136	4.279		-10.761		0.25
ATOM	1978	HB1		136				1.00	
					4.193		-10.365	1.00	0.27
ATOM	1979	HB2		136	3.302	-9.072	-11.064	1.00	0.26
MOTA	1980	ÇG	LEU	136	5.213	-8.709	-11.980	1.00	0.26
ATOM	1981	HG	LEU	136	5.312	-9.713	-12.368	1.00	0.29
MOTA	1982	CD1	LEU	136	4.624	-7.801	-13.063	1.00	0.29
ATOM	1983	HD11		136	3.546		-13.030		
ATOM	1984	HD12		136		-7.046	-13.030	1.00	1.06
					4.967	-8.126	-14.033	1.00	1.05
ATOM	1985	HD13		136	4.944	-6.784	-12.893	1.00	1.06
MOTA	1986	CD2		136	6.592		-11.578	1.00	0.32
MOTA	1987	HD21	LEU	136	6.485	-7.477	-10.762	1.00	1.05
ATOM	1988	HD22	LEU	136	7.046	-7 677	-12.422	1.00	1.09
MOTA	1989	HD23		136	7.220	-B 000	-11.269		
ATOM	1990	C	LEU	136		-0.336		1.00	0.97
		_			3.954	-9.556	-8.427	1.00	0.25
ATOM	1991	0	LEU	136	4.201	-8.761	-7.542	1.00	0.30
MOTA	1992	N	MET	137	2.924	-10.353	-8.357	1.00	0.28
MOTA	1993	HN	MET	137	2.744	-10.981	-9.087	1.00	0.31
MOTA	1994	CA	MET	137	2.016	-10.309	-7.177	1.00	
ATOM	1995	HA	MET	137	1.768				0.33
ATOM	1996					-9.283	-6.959	1.00	0.38
		CB	MET	137		-11.087	-7.494	1.00	0.42
MOTA	1997	HB1		137		-11.136	-6.615	1.00	0.57
MOTA	1998	HB2	MET	137	0.995	-12.089	-7.803	1.00	0.50
MOTA	1999	CG	MET	137	-0.035	-10.391	-8.625	1.00	0.58
MOTA	2000	HG1	MET	137	-0.909	-10.975	-8.875	1.00	
ATOM	2001	HG2		137					1.13
						-10.311	-9.494	1.00	1.22
ATOM	2002	SD	MET	137	-0.551	-8.729	-8.108	1.00	0.83
ATOM	2003	CE	MET	137	-2.048	-9.184	-7.194	1.00	0.39
MOTA	2004	HE1	MET	137	-2.231	-8.450	-6.426	1.00	1.14
ATOM	2005	HE2	MET	137	-1.927	-10.151	-6.741	1.00	1.07
ATOM	2006	HE3	MET	137	-2.885	-9.212	-7.872		
ATOM	2007	C	MET	137		-9.212		1.00	1.06
						-10.925	-5.951	1.00	0.27
MOTA	2008	0	MET	137		-11.287	-4.990	1.00	0.28
MOTA	2009	N	PHE	138	4.000	-11.042	-5.964	1.00	0.25
MOTA	2010	HN	PHE	138	4.514	-10.741	-6.743	1.00	0.28
ATOM	2011	ÇA	PHE	138		-11.628	-4.785	1.00	0.23
ATOM	2012	HA	PHE	138		-12.557			
ATOM	2013	CB					-4.534	1.00	0.26
			PHE	138	6.167	-11.877	-5.152	1.00	0.25
ATOM	2014	HB1		138		-10.945	-5.104	1.00	0.24
ATOM	2015		PHE	138	6.221	-12.270	-6.156	1.00	0.27
MOTA	2016	CG		138	6.790	-12.873	-4.194	1.00	0.28
ATOM	2017	CD1	PHE	138	6.295	-14.184	-4.113	1.00	0.32
ATOM	2018		PHE	138		-14.490	-4.731	1.00	
ATOM	2019	CD2	PHE	138	7 971	-12.486			0.33
ATOM	2020	מבי	PHE	138			-3.392	1.00	0.30
		AD2	PHE			-11.481		1.00	0.30
MOTA	2021		PHE	138	6.881	-15.100	-3.230	1.00	0.38
MOTA	2022		PHE	138	6.500	-16.109	-3.168	1.00	0.42
ATOM	2023	CE2	PHE	138		-13.404	-2.511	1.00	0.36
ATOM	2024	HE2	PHE	138		-13.104	-1.894	1.00	0.39
MOTA	2025	CZ	PHE	138		-14.710			
MOTA	2026	HZ					-2.430	1.00	0.39
			PHE	138		-15.417	-1.749	1.00	0.44
MOTA	2027	C	PHE	138	4.601	-10.615	-3.615	1.00	0.20
ATOM	2028	0	PHE	138	4.874	-9.447	-3.808	1.00	0.22
MOTA	2029	N	PRO	139		-11.019	-2.421	1.00	0.22
ATOM	2030	CA	PRO	139		-10.048			
ATOM	2031	HA	PRO	139			-1.291	1.00	0.25
ATOM					3.262	-9.340	-1.509	1.00	0.27
	2032	CB	PRO	139		-10.936	-0.127	1.00	0.31
ATOM	2033		PRO	139		-10.638	0.199	1.00	0.38
MOTA	2034	HB2	PRO	139		-10.835	0.691	1.00	0.42
MOTA	2035	CG	PRO	139		-12.392	-0.597	1.00	0.33
MOTA	2036	HG1		139		-12.812	-0.396		
ATOM	2037	HG2	PRO	139				1.00	0.41
ATOM	2038					-12.961	-0.074	1.00	0.42
		CD	PRO	139		-12.435	-2.102	1.00	0.27
MOTA	2039	HD2		139	4.661	-13.100	-2.318	1.00	0.28
MOTA	2040	HD1		139	2.946	-12.732	-2.637	1.00	0.30
MOTA	2041	C	DBU	120	E 227	-0.305	7 776		2.22

MOTA	2042	0	PRO	139	5.302	-8.351	-0.173	1.00	0.44
ATOM	2043	N	ILE	140	6.467	-9.726	-1.437		
ATOM	2044			_				1.00	0.24
		HN	ILE	140		-10.500	-2.038	1.00	0.37
ATOM	2045	CA	ILE	140	7.749	-9.031	-1.094	1.00	0.23
ATOM	2046	HA	ILE	140	7.572	-8.308	-0.312	1.00	0.24
ATOM	2047	CB	ILE	140					
						-10.054	-0.600	1.00	0.25
MOTA	2048	HB	ILE	140	8.978	-10.770	-1.379	1.00	0.25
ATOM	2049	CG1	ILE	140	8.207	-10.768	0.632	1.00	0.29
MOTA	2050	HG11	ILE	140		-11.196	0.384		
								1.00	0.32
MOTA		HG12	ILE	140		-10.055	1.434	1.00	0.33
MOTA	2052	CG2	ILE	140	10.070	-9.332	-0.214	1.00	0.26
MOTA	2053	HG21	ILE	140	9.850	-8.567	0.517	1.00	1.04
ATOM		HG22		140	10.505	-8.876	-1.090		
								1.00	1.06
MOTA		HG23		140		-10.040	0.207	1.00	1.04
MOTA	2056	CD1	ILE	140	9.156	-11.883	1.082	1.00	0.30
ATOM	2057	HD11	ILE	140		-12.250	0.236	1.00	1.08
ATOM		HD12		140		-12.691			
							1.511	1.00	0.98
MOTA	2059	HD13	ILE	140	9.838	-11.495	1.824	1.00	1.08
MOTA	2060	С	ILE	140	8.284	-8.301	-2.329	1.00	0.22
ATOM	2061	0	ILE	140	8.265	-8.817	-3.429	1.00	0.22
ATOM	2062								
		N	TYR	141	8.745	-7.092	2.150	1.00	0.21
MOTA	2063	HN	TYR	141	8.736	-6.696	-1.254	1.00	0.22
ATOM	2064	CA	TYR	141	9.265	-6.303	-3.304	1.00	0.21
MOTA	2065	HA	TYR	141	8.560	-6.348	-4.120	1.00	
MOTA	2066								0.20
		CB	TYR	141	9.444	-4.847	-2.865	1.00	0.21
MOTA	2067	HB1	TYR	141	10.050	-4.810	-1.972	1.00	0.22
MOTA	2068	HB2	TYR	141	8.476	-4.413	-2.661	1.00	0.22
ATOM	2069	CG	TYR	141	10.122				
						-4.066	-3.962	1.00	0.23
MOTA	2070	CDI	TYR	141	11.515	-4.104	-4.089	1.00	0.25
MOTA	2071	HD1	TYR	141	12.104	-4.697	-3.404	1.00	0.26
MOTA	2072	CD2	TYR	141	9.359	-3.298	-4.848	1.00	
ATOM	2073								0.24
		HD2	TYR	141	8.284	-3.268	-4.750	1.00	0.25
MOTA	2074	CE1	TYR	141	12.146	-3.376	-5.103	1.00	0.28
MOTA	2075	HE1	TYR	141	13.221	-3.405	-5.201	1.00	0.32
ATOM	2076	CE2	TYR	141	9.989				
						-2.569	-5.862	1.00	0.27
MOTA	2077	HE2	TYR	141	9.401	-1.975	-6.544	1.00	0.30
ATOM	2078	CZ	TYR	141	11.383	-2.608	-5.990	1.00	0.29
MOTA	2079	OH	TYR	141	12.005	-1.892	-6.991	1.00	0.33
ATOM	2080	НН		_					
			TYR	141	12.781	-2.385	-7.269	1.00	0.90
ATOM	2081	C	TYR	141	10.615	-6.864	-3.761	1.00	0.22
MOTA	2082	0	TYR	141	11.522	-7.050	-2.973	1.00	0.23
MOTA	2083	N	THR	142	10.750				
						-7.130	-5.035	1.00	0.22
MOTA	2084	HN	THR	142	10.002	-6.968	-5.648	1.00	0.22
MOTA	2085	CA	THR	142	12.035	-7.675	-5.563	1.00	0.24
ATOM	2086	HA	THR	142	12.835	-7.447	-4.874	1.00	
ATOM	2087	CB			11 015				0.25
			THR	142	11.917	-9.193	-5.723	1.00	0.25
MOTA	2088	HB	THR	142	11.645	-9.635	-4.777	1.00	0.26
MOTA	2089	OG1	THR	142	13.165	-9.720	-6.152	1.00	0.29
ATOM	2090	HG1	THR	142	13.274	-9.505			
ATOM	2091	CG2	THR	142			-7.081	1.00	0.97
					10.840	-9.517	-6.760	1.00	0.25
MOTA .		HG21	THR	142	10.577	-10.562	-6.691	1.00	1.04
MOTA	2093	HG22	THR	142	11.217	-9.304	-7.749	1.00	1.05
MOTA	2094	HG23	THR	142	9.965	-8.913			
ATOM	2095						-6.570	1.00	1.06
		Ç	THR	142	12.339	-7.040	-6.924	1.00	0.23
MOTA	2096	0	THR	142	11.454	-6.810	-7.724	1.00	0.23
ATOM	2097	N	TYR	143	· 13.586	-6.758	-7.195	1.00	0.25
MOTA	2098	HN	TYR	143	14.285	-6.955			
ATOM	2099	CA	TYR	143	13 040		-6.538	1.00	0.27
					13.948	-6.144	-8.506	1.00	0.26
MOTA	2100	HA	TYR	143	13.174	-5.452	-8.804	1.00	0.25
ATOM	2101	CB	TYR	143	15.277	-5.395	-8.370	1.00	0.29
ATOM	2102	HB1		143	16.072	-6.104			
ATOM				142			-8.190	1.00	0.33
	2103	HB2	TYR	143	15.217	-4.704	-7.542	1.00	0.30
ATOM	2104	CG	TYR	143	15.563	-4.633	-9.642	1.00	0.27
ATOM	2105	CD1	TYR	143	14.931	-3.406	-9.880	1.00	0.25
ATOM	2106	HD1		143					
					14.234	-3.008	-9.156	1.00	0.26
MOTA	2107		TYR	143	16.466	-5.148	-10.581	1.00	0.31
MOTA	2108	HD2		143	16.954		-10.398	1.00	0.35
MOTA	2109		TYR	143	15.201	-2 605	-11.055		
ATOM	2110	HE1				-2.075	-11.000	1.00	0.26
				143	14.713		-11.238	1.00	0.28
ATOM	2111	CE2	TYR	143	16.735	-4.436	-11.756	1.00	0.31
MOTA	2112	HE2	TYR	143	17.432	-4.833	-12.480	1.00	0.36
ATOM	2113	CZ	TYR	143	16.103	-3.210			
ATOM	2114	ОН					-11.994	1.00	0.28
			TYR	143	16.369	-2.509	-13.152	1.00	0.30
MOTA	2115	HH	TYR	143	17.068	-2.969	-13.624	1.00	0.95
ATOM	2116	С	TYR	143	14.080	-7.244	-9.563	1.00	0.27
ATOM	2117	Ō	TYR	143	14.552	-8.328			
ATOM	2118	Ŋ	THR	144		-6.328	-9.283	1.00	0.31
ALUM	~110				13 660	07E	10 220	1 00	^ ^^

MOTA	2119	HN	THR	144	13.277	-6 006	-10.972	1.00	0 22
MOTA	2120	CA	THR	144	13.753				0.32
							-11.847	1.00	0.32
MOTA	2121	HA	THR	144	14.479	-8.758	-11.573	1.00	0.35
MOTA	2122	CB	THR	144	12.385	-8.666	-12.031	1.00	0.37
MOTA	2123	HB	THR	144	11.922		-11.067	1.00	0.84
MOTA	2124	0G1	THR	144	12.549				
							-12.683	1.00	1.00
MOTA	2125	HG1	THR	144	13.280	-9.836	-13.301	1.00	1.42
MOTA	2126	CG2	THR	144	11.499	-7.757	-12.882	1.00	0.82
MOTA	2127	HG21	THR	144	10.461	-7 001	-12.699		
						-/.331	-12.099	1.00	1.51
MOTA	2128	HG22	THR	144	11.724	-7.911	-13.927	1.00	1.24
MOTA	2129	HG23	THR	144	11.687	-6.726	-12.622	1.00	1.49
MOTA	2130	С	THR	144	14.169		-13.165		
	2131							1.00	0.34
MOTA		0	THR	144	13.922		-13.392	1.00	0.32
MOTA	2132	N	GLY	145	14.789	-8.094	-14.043	1.00	0.43
ATOM	2133	HN	GLY	145	14.971	-9 037	-13.846	1.00	0.49
ATOM	2134	CA	GLY	145	15.205	7.510	15 350		
							-15.350	1.00	0.49
ATOM	2135		GLY	145	15.842	-8.207	-15.872	1.00	0.57
MOTA	2136	HA2	GLY	145	15.742	-6.587	-15.178	1.00	0.50
MOTA	2137	С	GLY	145	13.957		-16.191	1.00	
MOTA	2138	ō	GLY	145					0.47
					13.331	-8.138	-16.706	1.00	0.53
MOTA	2139	N	LYS	146	13.583	-5.990	-16.322	1.00	0.46
MOTA	2140	HN	LYS	146	14.097		-15.889	1.00	0.48
ATOM	2141	CA	LYS	146	12.367		-17.116		
ATOM	2142		LYS					1.00	0.49
	_	HA		146	11.578		-16.876	1.00	0.51
MOTA	2143	CB	LYS	146	11.911	-4.235	-16.764	1.00	0.52
MOTA	2144	HBl	LYS	146	10.973	-4.032	-17.254	1.00	0.58
MOTA	2145		LYS	146	12.657				
							-17.103	1.00	0.57
MOTA	2146	CG	LYS	146	11.744		-15.238	1.00	0.55
ATOM	2147	HG1	LYS	146	12.690	-3.853	-14.798	1.00	0.83
MOTA	2148	HG2	LYS	146	11.442	-6 000	-14.849		
MOTA	2149							1.00	1.14
		CD	LYS	146	10.684	-3.077	-14.854	1.00	1.23
ATOM	2150	HD1	LYS	146	10.308	~3.309	-13.871	1.00	1.78
MOTA	2151		LYS	146	9.865		-15.556	1.00	1.79
ATOM	2152	CE	LYS						
				146	11.298	-1.6/1	-14.828	1.00	2.01
MOTA	2153		LYS	146	11.615	-1.439	-13.822	1.00	2.47
MOTA	2154	HE2	LYS	146	10.556	-0.952	-15.143	1.00	2.39
ATOM	2155	NZ	LYS	146	12.468		-15.745		
MOTA	2156							1.00	2.91
			LYS	146	12.847		-15.750	1.00	3.39
ATOM	2157		LYS	146	12.170	-1.861	-16.707	1.00	3.28
MOTA	2158	HZ3	LYS	146	`13.205		-15.420	1.00	3.27
MOTA	2159	C	LYS	146	12.677				
							-18.613	1.00	0.59
MOTA	2160	0	LYS	146	11.845		-19.444	1.00	1.16
MOTA	2161	. И	SER	147	13.868	-6.131	-18.967	1.00	0.78
ATOM	2162	HN	SER	147	14.530	-6 366	-18.283		
MOTA	2163	CA	SER	147				1.00	1.26
					14.226		-20.413	1.00	0.87
MOTA	2164	HA	SER	147	14.141		-20.859	1.00	1.03
ATOM	2165	CB	SER	147	15.667	-6.709	-20.554	1.00	0.95
ATOM	2166	HB1	SER	147	15.798		-21.530		
ATOM	2167	HB2				-7.130	-21.530	1.00	1.42
				147	15.871	-7.445	-19.794	1.00	1.34
MOTA	2168	OG	SER	147	16.561	-5.616	-20.395	1.00	1.71
ATOM	2169	HG	SER	147	17.097		-21.190	1.00	2.16
MOTA	2170	C	SER	147	13.288		-21.138		
MOTA	2171							1.00	0.79
		0	SER	147	12.747	-6.865	-22.178	1.00	1.40
ATOM	2172	N	HIS	148	13.098	-8.366	-20.605	1.00	0.66
ATOM	2173	HN	HIS	148	13.551	-8.602	-19.768	1.00	1.10
ATOM	2174	CA	HIS	148	12.199	-0.360	-21.272		
ATOM	2175					-9.300	-21.2/2	1.00	0.65
		HA	HIS	148	11.629		-22.048	1.00	0.74
MOTA	2176	CB	HIS	148	13.041	-10.479		1.00	0.79
MOTA	2177	HB1	HIS	148	12.401	-11.312		1.00	
ATOM	2178		HIS	148					1.14
ATOM						-10.801	-21.174	1.00	1.30
	2179	CG	HIS	148	13.723	-9.980	-23.130	1.00	1.66
ATOM	2180		HIS	148	13.104	-9.116	-24.019	1.00	2.52
ATOM	2181	HD1	HIS	148	12.200	-0 747	-23.934		
ATOM	2182		HIS	148		-0./4/	-23.934	1.00	2.81
					14.969	-10.226	-23.652	1.00	2.62
MOTA	2183		HIS	148	15.715	-10.867	-23.206	1.00	3.00
MOTA	2184	CE1	HIS	148	13.970		-25.020	1.00	3.46
MOTA	2185		HIS	148	13.759				
ATOM						-0.233	-25.863	1.00	4.33
	2186		HIS	148	15.123	-9.528	-24.846	1.00	3.55
MOTA	2187	С	HIS	148	11.238	-9.971	-20.249	1.00	0.55
MOTA	2188	0	HIS	148		-11.064	-20 435		
ATOM	2189	N	PHE	149		-41.004	-20.433	1.00	0.60
					10.978		-19.167	1.00	0.57
ATOM	2190	HN	PHE	149	11.392	-8.417	-19.021	1.00	0.73
MOTA	2191	CA	PHE	149	10.060		-18.145	1.00	0.48
ATOM	2192	HA	PHE	149					
ATOM	2193					-10.849		1.00	0.51
		CB	PHE	149	10.022		-16.911	1.00	0.44
MOTA	2194	HB1	PHE	149	9.603	-8.008	-17.177	1.00	0.44
MOTA	2195	HB2	PHE	149	11.023		-16 530	1 00	0.49
							• • • • • • • • • • • • • • • • • • • •		· · · · -

ATOM ATOM	2196 2197	CG CD1		149 149	9.161 -9.615 -1 7.766 -9.507 -1	5.919	1.00	0.40 0.36
MOTA MOTA	2198 2199	HD1 CD2		149 149	7.305 -8.956 -1 9.757 -10.328 -1	.6.726 .4.804	1.00	0.38
MOTA	2200		PHE	149	10.832 -10.412 -1	4.750	1.00	0.42 0.48
MOTA	2201 2202	CE1		149		4.941	1.00	0.35
MOTA MOTA	2202	HE1 CE2	PHE	149 149		14.996 13.825	1.00	0.37
MOTA	2204		PHE	149	9.417 -11.482 -1	13.016	1.00	0.45
MOTA MOTA	2205 2206	CZ HZ	PHE PHE	149 149		13.894 13.140	1.00	0.37 0.38
MOTA	2207	С	PHE	149	8.641 -9.993 -1	18.706	1.00	0.43
MOTA MOTA	2208 2209	о И	PHE	149 150		19.217 18.575	1.00	0.45
MOTA	2210	HN	MET	150	8.523 -11.888 -1	18.133	1.00	0.50
MOTA MOTA	2211 2212	CA HA	MET MET	150 150	6.651 -11.357 -1 6.189 -10.400 -1	19.051	1.00	0.39 0.38
MOTA	2213	CB	MET	150	6.632 -12.207 -2		1.00	0.44
ATOM	2214 2215		MET	150		20.632	1.00	0.45
MOTA MOTA	2215	HB2 CG	MET MET	150 150	7.109 -13.157 -2 7.381 -11.477 -2	21.446	1.00	0.47
MOTA	2217	HG1		150	8.401 -11.831 -2	21.485	1.00	0.98
MOTA MOTA	2218 2219	HG2 SD	MET	150 150	7.376 -10.415 -2 6.571 -11.806 -2	21.253 23.033	1.00	0.86 1.32
MOTA	2220	CE	MET	150	7.378 -13.384 -2	23.393	1.00	2.23
MOTA MOTA	2221 2222	HE1 HE2	MET	150 150	7.326 -14.022 -2 8.411 -13.211 -2	22.521	1.00	2.66
MOTA	2223	HE3	MET	150	6.879 -13.861 -2	24.225	1.00	2.74
MOTA MOTA	2224 2225	CO	MET	150 150	5.877 -12.071 -1 6.435 -12.837 -1	17.943 17.183	1.00	0.32 0.32
MOTA	2226	N	LEU	151	4.605 -11.819 -1	17.827	1.00	0.32
MOTA MOTA	2227 2228	HN CA	LEU	151 151		18.437	1.00	0.30
MOTA	2229	HA	LEU	151	3.821 -12.478 -1 4.120 -12.064 -1	15.803	1.00	0.24
MOTA	2230	CB	LEU	151	2.327 -12.212 -1	16.966	1.00	0.24
MOTA MOTA	2231 2232		LEU	151 151	1.765 -12.626 -1 2.012 -12.680 -1	16.145 17.887	1.00	0.25 0.28
MOTA	2233	CG	LEU	151	2.061 -10.703 -1	17.04,7	1.00	0.28
MOTA MOTA	2234 2235	HG CD1	LEU	151 151	2.900 -10.208 -1 0.804 -10.457 -1	17.512	1.00	0.52 0.35
MOTA	2236	HD11	LEU	151	0.506 -9.424 -1	17.788	1.00	1.07
MOTA MOTA	2237 2238	HD12 HD13		151 151		17.526	1.00	1.02
MOTA	2239		LEU	151		18.917 15.638	1.00	1.17 0.46
MOTA MOTA		HD21 HD22	LEU	151 151	2.078 -9.084 -1 2.495 -10.650 -1		1.00	1.14
ATOM	2242	HD23		151	0.820 -10.284 -	14.941 15.345	1.00	1.16 1.11
MOTA MOTA	2243	C	LEU	151	4.076 -14.004 -:	16.794	1.00	0.24
ATOM	2244 2245	N 0	LEU PRO	151 152	3.879 -14.613 -1 4.504 -14.641 -1	17.826 15.711	1.00	0.28 0.22
MOTA	2246	CA	PRO	152	4.748 -16.112 -	15.751	1.00	0.23
MOTA MOTA	2247 2248	HA CB	PRO PRO	152 152	5.480 -16.354 -: 5.323 -16.404 -:	16.503 14.364	1.00 1.00	0.24
MOTA	2249	HB1	PRO	152	6.361 -16.686 -	14.453	1.00	0.29
MOTA MOTA	2250 2251	HB2 CG	PRO PRO	152 152	4.766 -17.208 -3 5.209 -15.141 -3		1.00	0.26 0.32
MOTA	2252	HG1	PRO	152	6.166 -14.917 -	13.061	1.00	0.44
MOTA MOTA	2253 2254	HG2 CD	PRO	152 152	4.473 -15.295 - 4.778 -13.976 -	12.730	1.00	0.41 0.25
MOTA	2255	HD2	PRO	152	3.886 -13.507 -	14.008	1.00	0.25
MOTA MOTA	2256 2257	HD1 C	PRO	152 152	5.581 -13.263 -	14.503	1.00	0.27
MOTA	2258	ŏ	PRO	152	3.462 -16.915 - 2.378 -16.371 -	16.038	1.00	0.21 0.20
MOTA	2259	N	ASP	153	3.582 -18.209 -	16.090	1.00	0.23
MOTA MOTA	2260 2261	HN CA	ASP ASP	153 153	4.468 -18.622 - 2.380 -19.063 -	16.031	1.00	0.25 0.23
MOTA	2262	HA	ASP	153	1.890 -18.772 -	17.221	1.00	0.23
ATOM ATOM	2263 2264	CB HB1	ASP ASP	153 153	2.813 -20.526 - 1.943 -21.163 -		1.00	0.25 0.26
MOTA	2265	HB2	ASP	153	3.470 -20.762 -	15.576	1.00	0.26
MOTA MOTA	2266 2267	CG	ASP ASP	153 153	3.550 -20.752 - 4.768 -20.687 -	17.722	1.00	0.27
MOTA	2268		ASP	153	2.884 -20.994 -	18.715	1.00	1.08 1.14
ATOM ATOM	2269 2270	CO	ASP ASP	153	1.409 -18.899 -	15.133	1.00	0.21
MOTA	2271	N	ASP	153 154	0.208 -18.858 - 1.919 -18.820 -		1.00	0.21 0.21
MOTA	2272	HM	ACD	154	2 801 -10 066 -		1 00	0.22

					4 440 44 484 44 44	
MOTA		CA	ASP	154		. 21
AYOM	2274	HA	ASP	154	0.431 -19.572 -12.641 1.00 0.	. 22
MOTA	2275	CB	ASP	154		. 23
ATOM		HB1	_	154		. 22
ATOM		HB2		154		
						. 25
ATOM		CG	ASP	154	0.975 -18.347 -10.267 1.00 0.	. 25
MOTA		OD1	ASP	154	1.276 -18.982 -9.269 1.00 1.	. 13
ATOM	2280	OD2	ASP	154	0.004 -17.613 -10.340 1.00 1.	. 07
ATOM	2281	С	ASP	154	0.102 -17.473 -12.943 1.00 0.	.19
ATOM	2282	ŏ	ASP	154		
						.19
MOTA	2283	N	ASP	155		.19
MOTA	2284	HN	ASP	155		.21
MOTA	2285	CA	ASP	155	-0.210 -15.140 -13.496 1.00 0.	.19
ATOM	2286	HA	ASP	155	-0.843 -15.011 -12.631 1.00 0	.20
MOTA	2287	CB	ASP	155		.21
MOTA	2288	HB1	ASP	155		.22
MOTA	2289	HB2		155		.22
ATOM	2290	CG	ASP	155		
	2291				1 111 111	.24
MOTA		OD1		155	2.355 -12.896 -12.327 1.00 1	.07
MOTA	2292	OD2		155		.14
MOTA	2293	С	ASP	155	-1.087 -15.300 -14.744 1.00 0	.19
ATOM	2294	0	ASP	155	-2.240 -14.918 -14.750 1.00 0	.19
ATOM	2295	N	VAL	156		.19
MOTA	2296	HN	VAL	156		.19
ATOM	2297	CA	VAL	156		
	2298				-1.372 -16.013 -17.041 1.00 0	.21
MOTA		AH	VAL	156		.22
ATOM	2299	CB	VAL	156		.23
MOTA	2300	HB	VAL	156	-0.034 -17.521 -17.776 1.00 0	.23
MOTA	2301	CG1	VAL	156		.27
ATOM	2302 H	G11	VAL	156		.00
ATOM			VAL	156		.05
ATOM	2304 H					
				156		.05
MOTA	2305	CG2		156		.26
MOTA	2306 H			156		.07
ATOM	2307 H	IG22	VAL	156	0.067 -14.856 -19.204 1.00 1	.05
ATOM	2308 F	IG23	VAL	156		.00
ATOM	2309	С	VAL	156		.20
ATOM	2310	0	VAL	156		.21
ATOM	2311	N	GLN	157		
ATOM	2312					.20
		HN	GLN	157		.20
ATOM	2313	CA	GLN	157		.22
MOTA	2314	HA	GLN	157		.24
MOTA	2315	CB	GLN	157	-2.995 -20:204 -15:117 1:00 0	.24
MOTA	2316	HB1	GLN	157		.26
ATOM	2317	HB2	GLN	157		.23
ATOM	2318	CG	GLN	157		.25
ATOM	2319		GLN	157		
ATOM	2320		GLN	157	-1.174 -20.686 -16.152 1.00 0	.94
						.87
ATOM	2321	CD	GLN	157		.19
ATOM	2322		GLN	157		.89
MOTA	2323	NE2		157	-1.646 -23.437 -16.364 1.00 1	.96
ATOM	2324 1	E21	GLN	157	-1.291 -23.203 -17.247 1.00 2	.18
MOTA	2325 F	E22	GLN	157		.65
ATOM	2326	C	GLN	157		.22
MOTA	2327	0	GLN	157		.24
ATOM	2328	N	GLY	158		
MOTA	2329	HN	GLY	158		.21
ATOM	2330					.20
		CA	GLY	158		.22
MOTA	2331	TAL	GLY	158	-4.380 -16.319 -12.232 1.00 0	.22
MOTA	2332	HA2		158	-5.667 -17.446 -12.646 1.00 0	.25
MOTA	2333	C	GLY	158		.20
MOTA	2334	0	GLY	158		.21
MOTA	2335	N	ILE	159		.18
ATOM	2336	HN	ILE	159		.18
ATOM	2337	CA	ILE			
	2338			159	-5.713 -13.593 -15.097 1.00 0	.19
MOTA		HA	ILE	159	-6.301 -13.054 -14.375 1.00 0	.20
MOTA	2339	CB	ILE	159	-4.679 <b>-12.648 -15.735 1.00 0</b>	.19
MOTA	2340	HB	ILE	159	-3.950 -12.367 -14.988 1.00 0	.20
ATOM	2341	CG1	ILE	159		.24
ATOM	2342 F			159		.26
MOTA		IG12		159		.28
ATOM	2344		ILE	159		
MOTA	2345 F	1621	TLE	159	-3.968 -13.361 -16.880 1.00 0	.21
ATOM	2345 F	1000	TITE		-2.998 -12.914 -17.036 1.00 1	.01
	2340 }	1022	TIE	159	-4.556 -13.274 -17.781 1.00 1	.01
MOTA	2347 F			159		.04
MOTA.	2348		ILE	159		.27
ATOM	2349 I	ID11	.ILE	159	-6.322 -9.644 -15 476 1 00 1	05

ATOM	2350	HD12	ILE	159	-4.644 -9.838 -14.978 1.00 1.	06
MOTA	2351	HD13	ILE	159	-5.893 -10.848 -14.265 1.00 1.	02
MOTA	2352	С	ILE	159		21
ATOM	2353		ILE	159		
						23
ATOM	2354		GLN	160		22
ATOM	2355	HN	GLN	160	-5.322 -15.538 -16.726 1.00 0.	21
ATOM	2356	CA	GLN	160	<b>-7.097 -15.763 -17.930 1.00 0.</b>	27
MOTA	2357		GLN	160		29
MOTA	2358		GLN	160		31
MOTA	2359	HB1	GLN	160	-6.999 -17.334 -19.389 1.00 0.	35
ATOM	2360	HB2	GLN	160		30
MOTA	2361		GLN	160		
						34
MOTA	2362	HG1		160		92
ATOM	2363	HG2	GLN	160	-5.799 -15.378 -20.290 1.00 0.	91
MOTA	2364	CD	GLN	160	-4.508 -17.087 -20.451 1.00 1.	11
ATOM	2365	OE1		160	-4.451 -18.248 -20.100 1.00 1.	88
MOTA	2366	NE2		160		. 83
ATOM	2367	HE21	GLN	160	-3.947 -15.767 -21.824 1.00 2.	.13
ATOM	2368	HE22	GLN	160		46
ATOM	2369	C	GLN	160		28
MOTA	2370	0	GLN	160	-9.386 -16.449 -17.779 1.00 0.	.31
MOTA	2371	N	SER	161	-8.086 -17.035 -16.117 1.00 0.	.27
MOTA	2372	HN	SER	161		.25
ATOM	2373	CA	SER	161		30
ATOM	2374					
		HA	SER	161		.34
ATOM	2375	CB	SER	161	-8.690 -18.427 -14.174 1.00 0.	. 33
MOTA	2376	HB1	SER	161	-7.861 -19.06714.444 1.00 O.	.35
ATOM	2377	HB2	SER	161		.36
					-5.470 *15.024 -15.741 1.00 U	
MOTA	2378	OG	SER	161		.33
MOTA	2379	HG	SER	161	-9.045 <b>-</b> 16.986 <b>-</b> 12.915 1.00 0.	. 94
ATOM	2380	С	SER	161	-10.267 -16.684 -15.019 1.00 0.	.30
ATOM	2381	0	SER	161		.35
ATOM	2382	Ň	LEU	162		
						. 27
MOTA	2383	HN	LEU	162		. 26
ATOM	2384	CA	LEU	162	-10.852 -14.413 -14.405 1.00 0.	. 29
MOTA	2385	HA	LEU	162		.33
ATOM	2386	CB	LEU	162		
						.28
MOTA	2387		LEU	162	-10.802 -12.509 -13.411 1.00 0	. 29
ATOM	2388	HB2	LEU	162	-9.256 -13.017 -14.086 1.00 0	. 27
MOTA	2389	CG	LEU	162		.30
ATOM	2390	HG	LEU	162		.30
MOTA	2391		LEU	162	-8.883 -12.918 -11.450 1.00 0	.33
MOTA	2392	HD11	LEU	162	-8.496 -13.370 -10.549 1.00 1	. 03
MOTA	2393	HD12	LEU	162	-9.490 -12.063 -11.191 1.00 1	.01
ATOM		HD13		162		.12
			-			
ATOM	2395		LEU	162		.33
MOTA	2396	HD21		162		.05
MOTA	2397	HD22	LEU	162	-11.812 -13.664 -11.697 1.00 1	.09
MOTA	2398	HD23	LEU	162		.01
MOTA	2399	C	LEU	162		.30
ATOM	2400	0	LEU	162		.36
ATOM	2401	N	TYR	163	-10.645 -13.300 -16.564 1.00 0	. 27
ATOM	2402	HN	TYR	163	-9.677 -13.404 -16.452 1.00 0	.26
ATOM	2403	CA	TYR	163		.31
ATOM	2404					
		HA	TYR	163		.33
MOTA	2405	CB	TYR	163		.29
MOTA	2406	HBI	TYR	163	-10.562 -11.112 -19.170 1.00 0	.32
ATOM	2407	HB2	TYR	163	-9.234 -11.952 -18.371 1.00 0	.29
MOTA	2408	CG	TYR	163		.25
MOTA	2409		TYR	163		
						.23
MOTA	2410		TYR	163		. 23
MOTA	2411	CD2	TYR	163	-11.042 -9.357 -17.258 1.00 0	.27
ATOM	2412	HD2	TYR	163		.30
MOTA	2413	CEL		163		.24
MOTA	2414	HE1		163		.25
MOTA	2415	CE2		163	-10.984 -8.348 -16.289 1.00 0	.27
MOTA	2416	HE2	TYR	163		.30
ATOM	2417	CZ	TYR	163		.27
ATOM	2418					
		OH	TYR	163		.31
MOTA	2419	HH	TYR	163	-10.344 -7.782 -13.481 1.00 0	.99
ATOM	2420	C	TYR	163		.37
ATOM	2421	0	TYR	163		.43
ATOM	2422	Ŋ	GLY	164		
						.38
ATOM	2423	HN	GLY	164		.35
ATOM	2424	CA	GLY	164		.47
MOTA	2425	HA1	GLY	164		.53
D TVOM	2425	UND		164	14 140 15 074 10 205 1 00 0	

MOTA	2427	C	GLY	164	-9.735	-15.902	-20.648	1.00	0.55
ATOM	2428	0	GLY	164	-9.761	-15.580	-21.819	1.00	1.01
TER	2429		GLY	164					
HETATM		ZN	ZN	166	-0.218	-6.515	-2.613	1.00	0.24
HETATM		ZN	ZN	167	-3.506	6.833	-0.714	1.00	0.97
HETATM	2432	CA	CA	168	6.060	3.350	3.030	1.00	0.23
HETATM	2433	Cl	WAY	169	2.180	-4.315	1.627	0.00	0.30
HETATM	2434	C2	WAY	169	0.865	-4.629	1.215	0.00	0.33
HETATM	2435	1CE1	WAY	169	-0.170	-4.517	2.143	0.00	0.38
HETATM	2436	1CZ	WAY	169	0.074	-4.157	3.457	0.00	0.40
HETATM	2437	1CE2	WAY	169	1.355	-3.807	3.841	0.00	0.38
HETATM	2438	C6	WAY	169	2.395	-3.805	2.922	0.00	0.33
HETATM	2439	1HE1	WAY	169	-1.190	~4.713	1.839	0.00	0.42
HETATM			WAY	169	-0.734	-4.151	4.173	0.00	0.45
HETATM		1HE2		169	1.535	-3.534	4.872	0.00	0.42
HETATM			WAY	169	0.444	-5.080	-0.136	0.00	0.36
HETATM			WAY	169	0.467	-6.264	-0.463	0.00	0.58
HETATM		N12		169	-0.019	-4.195	-1.032	0.00	
HETATM			WAY	169	-0.045	-4.608	-2.371	-	0.61
HETATM			WAY	169	-0.357	-3.297	-0.743	0.00	0.68
HETATM			WAY	169	-0.953	-4.727		0.00	0.88
HETATM		1CH1		169	3.728		-2.645	0.00	1.13
HETATM		1HH1		169	3.702	-3.247	3.360	0.00	0.37
HETATM		1HH2		169	4.519	-2.162	3.422	0.00	1.07
HETATM		1HH3		169	4.013	-3.516	2.664	0.00	1.06
HETATM	_		WAY	169	3.274	-3.623	4.339	0.00	1.11
HETATM			WAY	169	3.865	-4.485 -3.175	0.819	0.00	0.29
HETATM			WAY	169	3.882	-5.812	0.021	0.00	0.25
HETATM		2CE1		169			0.684	0.00	0.32
HETATM			WAY	169	7.334	-6.241	2.178	0.00	1.09
HETATM			WAY	169	6.971	-6.520	3.488	0.00	0.53
HETATM		2CD2			5.697	-6.659	3.876	0.00	1.47
HETATM			WAY	169 169	4.747	-6.451	2.954	0.00	1.37
HETATM		2CD1			5.010	-6.084	1.640	0.00	0.36
HETATM		2HE1		169	6.338	-5.982	1.250	0.00	1.14
HETATM			WAY	169 169	8.374	-6.224	1.881	0.00	1.94
HETATM		2HD2		169	7.752	-6.630	4.227	0.00	0.61
HETATM		2HD1		169	3.708	-6.570	3.227	0.00	2.23
HETATM		2HB1			6.599	-5.706	0.239	0.00	2.05
HETATM		2HB2		169 169	4.245	-5.905	-0.339	0.00	0.31
HETATM			WAY		3.095	-6.552	0.832	0.00	0.34
HETATM		3CD1		169	4.187	-3.617	-1.665	0.00	0.23
HETATM		3CE1		169 169	3:310	-3.216	-2.661	0.00	0.25
HETATM			WAY	169	3.622	-3.465	-3.992	0.00	0:27
HETATM		3CE2		169	4.769	-4.183	-4.326	0.00	0.24
HETATM		3CD2		169	5.602	-4.644	-3.308	0.00	0.23
HETATM		3HD1		169	5.315	-4.359	-1.979	0.00	0.23
HETATM		3HE1			2.392	-2.714	-2.389	0.00	0.29
HETATM		3HE2		169 169	2.961	-3.091	-4.758	0.00	0.31
HETATM		3HD2		169	6.481	-5.228	-3.535	0.00	0.26
HETATM			WAY	169	5.959	-4.707	-1.184	0.00	0.27
HETATM			WAY	169	5.078	-4.439	-5.664	0.00	0.27
HETATM		3HH1		169	6.245	-5.202	-5.904	0.00	0.28
HETATM		3HH2		169	6.379	-5.372	-6.973	0.00	0.31
HETATM		3HH3		169	6.178	-6.172	-5.407	0.00	0.28
HETATM			WAY	169	7.127	-4.683	-5.526	0.00	0.29
HETATM			WAY	169	5.123	-2.847	0.614	0.00	0.27
END		031	*****	103	2.834	-2.186	0.004	0.00	0.25

	Atom Type	Res.		x	Y	z	Occ. B	MOL.
MOTA	1 CB	THR	7	73.468	27.410	6.079	1.00 42.70	A_13
MOTA	2 OG1	THR	7	72.149	27.911	6.358	1.00 37.82	A_13
ATOM ATOM	4 CG2 5 C	THR THR	7 7	73.843 75.936	26.297 28.076	7.068 6.227	1.00 25.79 1.00 28.29	A_13
ATOM	6 0	THR	ż	76.497	28.090	7.332	1.00 28.29 1.00 22.94	A_13 A_13
MOTA	9 N	THR	7	74.360	29.396	4.862	1.00 20.25	A_13
ATOM	11 CA	THR	7	74.501	28.593	6.099	1.00 21.49	A_13
MOTA MOTA	12 N 14 CA	LEU LEU	8 8	76.547 77.915	27.691 27.150	5.099 5.105	1.00 32.90 1.00 31.85	A_13
ATOM	15 GB	LEU	8	77.952	25.759	4.438	1.00 31.85 1.00 21.38	A_13 A_13
MOTA	16 CG	LEU	8	78.016	25.576	2.910	1.00 29.31	A_13
ATOM		LEU LEU	8 8	79.463	25.509	2.425	1.00 16.78	A_13
ATOM ATOM	19 C	LEU	8	77.334 78.956	24.292 28.070	2.527 4.465	1.00 23.37 1.00 24.01	A_13 A_13
ATOM	20 O	LEU	8	78.835	28.415	3.293	1.00 26.18	A_13
MOTA	21 N	LYS	9	79.980	28.424	5.251	1.00 36.26	A_13
ATOM ATOM	23 CA 24 CB	LYS LYS	9 9	81.106 82.438	29.298 28.521	4.867 4.977	1.00 23.24 1.00 25.52	A_13
ATOM	25 CG	LYS	é	82.767	27.570	3.815	1.00 25.52	A_13 A_13
MOTA	26 CD	LYS	9	83.661	28.243	2.753	1.00 31.69	A_13
ATOM ATOM	27 CE 28 NZ	LYS LYS	9 9	83.451	27.688	1.323 0.797	1.00 25.30	A_13
ATOM	32 C	LYS	9	82.056 81.042	27.938 30.073	3.526	1.00 20.65 1.00 31.41	A_13 A_13
MOTA	33 O	LYS	9	80.764	29.505	2.466	1.00 22.31	A_13
MOTA	34 N	TRP	10	81.327	31.372	3.573	1.00 15.84	A_13
ATOM ATOM	36 CA 37 CB	TRP TRP	10 10	81.312 81.636	32.172 33.620	2.361 2.680	1.00 10.58 1.00 21.39	A_13 A_13
MOTA	38 CG	TRP	10	BO.529	34.337	3.343	1.00 22.84	A_13
MOTA	39 CD2	TRP	10	79.479	35.074	2.697	1.00 20.41	A_13
MOTA MOTA	40 CE2 41 CE3	TRP TRP	10 10	78.676 79.142	35.631 35.320	3.718 1.357	1.00 24.50	A_13
ATOM	42 CD1		10	80.327	34.469	4.682	1.00 13.29 1.00 13.40	A_13 A_13
ATOM	43 NE1	TRP	10	79.220	35.253	4.919	1.00 18.40	A_13
ATOM	45 CZ2	TRP	10	77.550	36.418	3.442	1.00 12.63	A_13
MOTA MOTA	46 CZ3 47 CH2		10 10	78.021 77.242	36.105 36.641	1.083 2.120	1.00 19.89 1.00 13.62	A_13 A_13
MOTA	48 C	TRP	10	82.377	31.594	1.455	1.00 22.95	A_13
MOTA.	49 0	TRP	10	83.450	31.221	1.920	1.00 16.28	A 13
MOTA MOTA	50 N 52 CA	SER SER	11 11	82.087 83.017	31.533 30.975	0.167 -0.801	1.00 14.81 1.00 19.50	A_13 A_13
MOTA	53 CB	SER	11	82.282	30.596	-2.086	1.00 24.36	A_13
MOTA	54 OG	SER	11	81.605	29.353	-1.958	1.00 40.49	A 13
MOTA MOTA	56 C 57 O	SER SER	11 11	84.190 85.132	31.867 31.423	-1.134 -1.779	1.00 16.53	A_13
MOTA	58 N	LYS	12	84.153	33.113	-0.686	1.00 23.48 1.00 12.50	A_13 A_13
ATOM	60 CA	LYS	12	85.232	34.057	-0.961	1.00 17.05	A_13
MOTA MOTA	61 CB 62 CG	LYS LYS	12	84.741	35.168	-1.891	1.00 17.32	A_13
ATOM	63 CD	LYS	12 12	83.526 82.788	35.898 36.644	-1.350 -2.446	1.00 18.49	A_13 A_13
ATOM	64 CE	LYS	12	81.534	37.282	-1.888	1.00 18.44	A_13
ATOM	65 NZ	LYS	12	80.805	38.094	-2.895	1.00 16.65	A_13
atom atom	69 C 70 O	Lys Lys	12 12	85.687 84.946	34.662 34.637	0.344 1.319	1.00 11.16 1.00 12.63	A_13 A_13
ATOM	71 N	MET	13	86.915	35.185	0.355	1.00 15.52	A_13
MOTA	73 CA 74 CB	MET	13	87.516	35.801	1.537	1.00 11.04	A_13
MOTA MOTA	74 CB 75 CG	MET MET	13 13	89.028 89.431	35.547 34.082	1.565 1.707	1.00 16.57 1.00 20.92	A_13 A_13
MOTA	76 SD	MET	13	88.905	33.235	3.227	1.00 20.10	A_13
ATOM	77 CE	MET	13	87.486	32.313	2.604	1.00 16.29	A_13
MOTA MOTA	78 C 79 O	MET	13 13	87.258 87.247	37.296 37.916	1.572 2.634	1.00 13.23 1.00 22.80	A_13 A_13
ATOM	80 N	ASN	14	87.111	37.875	0.389	1.00 15.02	A_13
MOTA	82 CA	ASN	14	86.853	39.294	0.241	1.00 33.02	A_13
MOTA MOTA	83 CB 84 CG	asn Asn	14 14	87.445 88.925	39.801	-1.082	1.00 19.42	A_13
ATOM		. Asn	14	89.343	39.482 38.341	-1.217 -1.031	1.00 30.32 1.00 30.12	A_13 A_13
MOTA	86 ND2	2 ASN	14	89.723	40.489	-1.549	1.00 28.22	A_13
MOTA	89 C	ASN	14	85.337	39.482	0.277	1.00 27.58	A_13
MOTA MOTA	90 O 91 N	asn Leu	14 15	84.606 84.868	38.935 40.212	-0.568 1.287	1.00 28.01 1.00 19.06	A_13 A_13
ATOM	93 CA	LEU	15	83.444	40.450	1.459	1.00 20.03	Ã_13
MOTA	94 CB	LEU	15	82.930	39.690	2.691	1.00 19.55	A_13
MOTA MOTA	95 CG 96 CD	LEU L LEU	15 15	83.027 83.216	38.166 37.555	2.593 3.962	1.00 19.02 1.00 17.48	A_13 A_13
ATOM		LEU	15	81.799	37.604	1.903	1.00 23.43	A_13
ATOM	98 C	LEU	15	83.161	41.928	1.609	1.00 19.52	A_13
MOTA	99 O	LEU	15	83.980	42.676	2.130	1.00 15.98	A_13

FIG. 5

MOTA	100	N	THR	16	81.983	42.343	1.162	1.00 21.22	2 13
ATOM	102	CA	THR	16	81.578	43.736	1.252	1.00 10.00	A_13
ATOM	103	CB	THR	16	81.194	44.257	-0.109	1.00 10.00	A_13
MOTA	104	OG1		16	80.225	43.370	-0.681		A_13
ATOM	106	CG2	THR	16				1.00 22.43	A_13
ATOM	107				82.427	44.383	-1.009	1.00 15.42	A_13
		Č	THR	16	80.368	43.869	2.184	1.00 14.48	A_13
ATOM	108	0	THR	16	79.647	42.897	2.445	1.00 15.74	A_13
MOTA	109	И	TYR	17	80.176	45.065	2.716	1.00 15.89	A_13
ATOM	111	CA	TYR	17	79.064	45.340	3.604	1.00 13.19	A_13
ATOM	112	CB	TYR	17	79.480	45.195	5.067	1.00 21.42	A_13
MOTA	113	CG	TYR	17	80.448	46.236	5.580	1.00 26.23	A_13
ATOM	114	CD1	TYR	17	81.824	46.081	5.412	1.00 16.37	A_13
ATOM	115	CE1	TYR	17	82.724	46.981	5.988	1.00 12.90	A_13
ATOM	116		TYR	17	79.990	47.329	6.331	1.00 17.15	N_13
MOTA	117	CE2		· 17	80.880	48.235	6.912	1.00 24.15	A_13
ATOM	118	CZ	TYR	17	82.244				A_13
ATOM	119	OH	TYR	17	02.244	48.057	6.743	1.00 23.38	A_13
ATOM	121				83.121	48.942	7.343	1.00 19.47	A_13
		C	TYR	17	78.573	46.740	3.343	1.00 10.00	A_13
ATOM	122	0	TYR	17	79.298	47.559	2.782	1.00 19.27	A_13
ATOM	123	N	ARG	18	77.349	47.019	3.762	1.00 18.52	A_13
ATOM	125	CA	ARG	18	76.762	48.332	3.577	1.00 10.00	A_13
MOTA	126	CB	ARG	18	75.970	48.363	2.274	1.00 10.00	A_13
MOTA	127	CG	ARG	18	75.134	49.619	2.094	1.00 14.01	A_13
MOTA	128	CD	ARG	18	74.266	49.524	0.846	1.00 13.91	A_13
MOTA	129	NE	ARG	18	73.298	50.615	0.782	1.00 13.55	A_13
ATOM	131	CZ	ARG	18	72.165	50.571	0.092	1.00 10.00	A_13
ATOM	132	NH1	ARG	18	71.855	49.488	-0.602	1.00 14.30	A_13
MOTA	135	NH2	ARG	18	71.331	51.604	0.125	1.00 28.79	
ATOM	138	С	ARG	18	75.842	48.640	4.741	1.00 10.65	A_13
ATOM	139	ō.	ARG	18	75.037	47.796	5.141		A_13
ATOM	140	N	ILE	19				1.00 12.86	A_13
ATOM	142	CA	ILE		76.014	49.814	5.332	1.00 25.54	A_13
ATOM				19	75.169	50.265	6.436	1.00 24.52	A_13
	143	CB	ILE	19	75.944	51.236	7.350	1.00 18.37	A_13
ATOM	144		ILE	19	75.034	51.765	8.485	1.00 13.87	A_13
MOTA	145		ILE	19	77.204	50.545	7.888	1.00 27.67	A_13
MOTA	146		ILE	19	78.203	51.501	8.557	1.00 22.81	A_13
MOTA	147	C	ILE	19	74.062	51.027	5.698	1.00 21.11	A_13
ATOM	148	0	ILE	19	74.261	52.179	5.300	1.00 10.00	A_13
MOTA	149	N	VAL	20	72.916	50.378	5.487	1.00 19.76	Ā_13
MOTA	151	CA	VAL	20	71.829	51.014	4.735	1.00 18.20	
ATOM	152	CB	VAL	20	70.774	49.983	4.193		A_13
ATOM	153		VAL	20	71.384			1.00 15.42	A_13
ATOM	154		VAL			48.570	4.088	1.00 10.00	A_13
ATOM	155			20	69.496	50.030	4.992	1.00 18.62	A_13
ATOM		C	VAL	20	71.175	52.206	5.443	1.00 11.67	A_13
	156	0	VAL	20	70.652	53.110	4.798	1.00 18.36	A_13
ATOM	157	N	ASN	21	71.153	52.187	6.773	1.00 10.94	A_13
ATOM	159	CA	ASN	21	70.609	53.316	7.544	1.00 11.99	A_13
ATOM	160	CB	asn	21	69.078	53.307	7.675	1.00 10.00	A_13
MOTA	161	CG	ASN	21	68.533	51.978	8.107	1.00 14.93	A_13
ATOM	162	OD1	asn	21	67.627	51.449	7.486	1.00 21.54	A_13
ATOM	163	ND2	ASN	21	69.105	51.408	9.148	1.00 10.00	A_13
ATOM	166	С	ASN	21	71.291	53.382	8.897	1.00 18.90	A_13
ATOM	167	0	ASN	21	72.006	52.447	9.283	1.00 12.49	A_13
ATOM	168	N	TYR	22	71.053	54.471		1.00 17.47	A_13
ATOM	170	CA	TYR	22	71.681	54.708	10.910	1.00 24.85	A_13
MOTA	171	СВ	TYR	22	72.556	55.954	10.818	1.00 13.52	A_13
ATOM	172	CG	TYR	22	73.791	55.748	9.991		A_13
MOTA	173		TYR	22	75.033			1.00 10.00	A_13
ATOM	174	CE1		22		55.600	10.598	1.00 14.05	A_13
ATOM	175				76.180	55.370	9.841	1.00 13.69	A_13
ATOM		CD2		22	73.717	55.663	8.608	1.00 10.00	A_13
	176	CE2	TYR	22	74.848	55.432	7.847	1.00 17.10	A_13
ATOM	177	CZ	TYR	22	76.077	55.288	8.476	1.00 14.43	A_13
MOTA	178	OH	TYR	22	77.204	55.072	7.737	1.00 10.00	A_13
ATOM	180	C	TYR	22	70.726	54.862	12.076	1.00 25.95	A_13
MOTA	181	0	TYR	22	69.593	55.311	11.916	1.00.10.00	A_13
ATOM	182	N	THR	23	71.187	54.483	13.259	1.00 20.30	N 12
MOTA	184	CA	THR	23	70.367	54.606	14.450	1.00 20.30	A_13
ATOM	185	CB	THR	23	70.821		15 504		A_13
ATOM	186		THR	23		53.635	15.584	1.00 10.90	A_13
ATOM	188				70.136	53.968	16.792	1.00 10.00	A_13
ATOM			THR	23	72.328	53.752	15.852	1.00 16.51	A_13
	189	Ç	THR	23	70.459	56.038	14.959	1.00 18.14	A_13
ATOM	190	0	THR	23	71.360	56.785	14.575	1.00 10.00	A_13
MOTA	191	N	PRO	24	69.433	56.487	15.691	1.00 12.76	A_13
ATOM	192	CD	PRO	24	68.061	55.950	15.716	1.00 15.26	A_13
MOTA	193	CA	PRO	24	69.453	57.844	16.232	1.00 22.70	A_13
MOTA	194	CB	PRO	24	67.985	58.086	16.585	1.00 28.52	A_13
MOTA	195	CG	PRO	24	67.448	56.706	16.841	1.00 15.78	A_13

ATOM	196	С	PRO	24	70.346	57.945	17.475	1.00 24.52	
MOTA	197	ŏ	PRO	24	70.790	59.040	17.831	1.00 24.52	A_13 A_13
ATOM	198	N	ASP	25	70.614	56.797	18.105	1.00 11.82	A_13
MOTA	200	CA'	ASP	25	71.416	56.721	19.336	1.00 12.31	A_13
MOTA	201	CB	ASP	25	71.339	55.317	19.917	1.00 25.26	A_13
ATOM	202	CG	ASP	25	69.927	54.782	19.977	1.00 10.00	A_13
MOTA	203	OD1		25	69.783	53.567	20.159	1.00 20.90	A_13
MOTA	204	OD2	ASP	25	68.960	55.558	19.841	1.00 18.45	A_13
MOTA	205	С	ASP	25	72.891	57.113	19.286	1.00 14.34	A_13
MOTA	206	0	ASP	25	73.449	57.511	20.301	1.00 11.77	A_13
MOTA	207	N	MET	26	73.546	56.873	18.157	1.00 20.78	A_13
MOTA	209	CA.	MET	26	74.960	57.208	18.010	1.00 20.03	A_13
MOTA	210	CB	MET	26	75.791	55.928	17.916	1.00 13.86	A_13
MOTA	211	CG	MET	26	75.966	55.181	19.231	1.00 19.00	A_13
MOTA	212	SD	MET	26	76.043	53.404	18.941	1.00 14.67	A_13
ATOM	213	CE	MET	26	77.737	53.223	18.385	1.00 19.74	A_13
ATOM	214	C	MET	26	75.157	58.047	16.754	1.00 13.32	A_13
ATOM	215	0	MET	26	74.274	58.086	15.900	1.00 16.81	A_13
MOTA	216	N	THR	27	76.285	58.749	16.656	1.00 10.29	A_13
MOTA MOTA	218 219	CA CB	THR THR	27 27	76.568	59.564	15.470	1.00 17.00	A_13
ATOM	220	OG1	THR	27	77.710 78.969	60.596	15.700	1.00 11.79	A_13
ATOM	222	CG2	THR	27	77.519	59.921 61.342	15.729 17.020	1.00 23.77 1.00 21.98	A_13
MOTA	223	C	THR	27	76.996	58.634	14.347	1.00 21.38	A_13 A_13
ATOM	224	ō	THR	27	77.411	57.500	14.608	1.00 11.05	A_13 A_13
ATOM	225	N	HIS	28	76.972	59.124	13.113	1.00 10.00	A_13
ATOM	227	CA	HIS	28	77.362	58.300	11.980	1.00 10.96	A_13
ATOM	228	CB	HIS	28	77.240	59.071	10.657	1.00 16.07	A_13
MOTA	229	CG	HIS	28	75.829	59.382	10.264	1.00 15.53	A_13
MOTA	230	CD2	HIS	28	74.707	59.531	11.016	1.00 21.47	A_13
MOTA	231	ND1	HIS	28	75.440	59.597	8.959	1.00 30.32	A_13
ATOM	233	CE1	HIS	28	74.149	59.868	8.920	1.00 19.38	A_13
MOTA	234	NE2	HIS	28	73.680	59.833	10.160	1.00 29.43	A_13
MOTA	236	С	HIS	28	78.769	57.735	12.151	1.00 14.80	A_13
MOTA	237	0	HIS	28	79.005	56.568	11.851	1.00 28.24	A_13
ATOM	238	N	SER	29	79.703	58.548	12.634	1.00 14.00	A_13
ATOM	240	CA	SER	29	81.068	58.070	12.854	1.00 19.57	A_13
ATOM	241	CB	SER	29	82.001	59.219	13.242	1.00 17.84	A_13
ATOM	242	OG	SER	29	82.383	59.936	12.084	1.00 28.25	A_13
MOTA	244	C	SER	29	81.134	56.983	13.917	1.00 15.23	A_13
MOTA	245	0	SER	29	81.818	55.973	13.733	1.00 13.73	A_13
MOTA	246	N	GLU	30	80.428	57.182	15.027	1.00 27.71	A_13
MOTA MOTA	248 249	CA	GLU	30	80.430	56.186	16.100	1.00 23.60	A_13
ATOM	250	CB	GLU	30	79.571	56.635	17.289	1.00 21.72	A_13
ATOM	251	CD	GLU	30 30	80.048	57.913	17.973	1.00 24.07	A_13
ATOM	252	OE1		30	79.205 79.784	58.279 58.660	19.185	1.00 21.06	A_13
ATOM	253	OE2		30	77.963	58.185	20.218 19.119	1.00 46.95 1.00 18.27	A_13
ATOM	254	c	GLU	30	79.895	54.877	15.553	1.00 18.27	A_13 A_13
MOTA	255	ō	GLU	30	80.456	53.809	15.815	1.00 13.06	A_13
MOTA	256	N	VAL	31	78.839	54.970	14.746	1.00 16.23	A_13
ATOM	258	CA	VAL	31	78.225	53.781	14.146	1.00 22.33	A_13
MOTA	259	CB	VAL	31	76.899	54.135	13.390	1.00 23.53	A_13
ATOM	260	CG1	VAL	31	76.384	52.920	12.628	1.00 14.39	A_13
MOTA	261	CG2	VAL	31	75.829	54.587	14.377	1.00 10.00	A_13
MOTA	262	C	VAL	31	79.208	53.040	13.216	1.00 20.29	A_13
MOTA	263	0	VAL	31	79.330	51.814	13.282	1.00 14.02	A_13
MOTA	264	N	GLU	32	79.913	53.790	12.370	1.00 23.94	A_13
ATOM	266	CA	GLU	32	80.887	53.219	11.446	1.00 10.18	A_13
MOTA	267	CB	GLU	32	81.406	54.285	10.502	1.00 16.50	A_13
MOTA	268	CG	GLU	32	80.424	54.605	9.427	1.00 20.84	A_13
ATOM ATOM	269	CD	GLU	32	80.330	56.080	9.155	1.00 22.31	A_13
ATOM	270		GLU	32	79.285	56.509	8.639	1.00 29.39	A_13
ATOM	271 272		GLU	32	81.294	56.812	9.458	1.00 22.01	A_13
ATOM	273	C	GLU GLU	32	82.056	52.565	12.137	1.00 18.93	A_13
ATOM	274	O N	LYS	32 33	82.474	51.470	11.753	1.00 24.42	A_13
ATOM	276	N CA	LYS	33 33	82.610	53.241	13.139	1.00 19.78	A_13
ATOM	277	CB	LYS	33	83.726 84.340	52.661	13.873	1.00 28.68	A_13
ATOM	278	CG	LYS	33	85.016	53.681	14.837	1.00 18.54	A_13
ATOM	279	CD	LYS	33	86.135	54.855 54.425	14.135	1.00 31.19	A_13
ATOM	280	CE	LYS	33	85.600	53.972	13.148 11.785	1.00 40.31	A_13
ATOM	281	NZ	LYS	33	86.646	53.779	10.773	1.00 21.99	A_13 ·
ATOM	285	c	LYS	33	83.242	51.407	14.594	1.00 33.20 1.00 12.66	A_13 A_13
ATOM	286	ŏ	LYS	33	83.892	50.361	14.552	1.00 12.66	A_13 A_13
ATOM	287	N	ALA	34	82.036	51.481	15.148	1.00 19.34	A_13
ATOM	289	CA	ALA	34	81.453	50.344	15.843	1.00 10.00	A_13

ATOM	290	CB	ALA	34	80.040	50.651	16.279	1.00 18.59	A_13
ATOM	291	C	ALA	34	81.468	49.119	14.940	1.00 13.45	A_13
ATOM ATOM	292	0	ALA	34	82.067	48.095	15.284	1.00 15.90	A_13
ATOM	293	N	PHE	35	80.857	49.234	13.766	1.00 19.57	A_13
ATOM	295 296	CA CB	PHE	35	80.802	48.112	12.812	1.00 26.77	A_13
ATOM	297	_	PHE	35	79.837	48.423	11.660	1.00 17.34	A_13
ATOM	298	CG	PHE PHE	35	78.390	48.477	12.077	1.00 30.55	A_13
ATOM	299			35	77.838	47.464	12.863	1.00 26.58	A_13
ATOM	300		PHE	35	77.570	49.512	11.653	1.00 10.00	A_13
MOTA	301		PHE PHE	35	76.494	47.485	13.212	1.00 12.45	A_13
ATOM	302	CZ	PHE	35 35	76.224	49.538	12.002	1.00 17.92	A_13
ATOM	303	C	PHE	35	75.684 82.170	48.525	12.777	1.00 13.29	A_13
ATOM	304	Ö	PHE	35		47.754	12.236	1.00 11.31	A_13
ATOM	305	N	LYS	36	82.493 82.962	46.573 48.778	12.034	1.00 11.37	A_13
ATOM	307	CA	LYS	36	84.293	48.573	11.945 11.400	1.00 17.06	A_13
ATOM	308	СВ	LYS	36	84.991	49.922	11.208	1.00 17.41 1.00 11.20	A_13
ATOM	309	CG	LYS	36	86.282	49.792	10.439	1.00 11.20	A_13
ATOM	310	CD	LYS	36	87.246	50.917	10.738	1.00 24.52	A_13 A_13
ATOM	311	CE	LYS	36	88.542	50.703	9.978	1.00 24.52	A_13 A_13
ATOM	312	NZ	LYS	36	88.264	50.536	8.514	1.00 23.69	A_13 A_13
MOTA	316	С	LYS	36	85.122	47.685	12.345	1.00 16.09	A_13
MOTA	317	0	LYS	36	85.701	46.686	11.938	1.00 21.50	A_13
MOTA	318	N	LYS	37	85.173	48.057	13.613	1.00 12.42	Ä_13
MOTA	320	CA	LYS	37	85.926	47.303	14.591	1.00 12.36	A_13
MOTA	321	CB	LYS	37	85.953	48.066	15.917	1.00 13.65	A_13
ATOM	322	CG	LYS	37	86.744	47.374	17.028	1.00 13.38	A_13
ATOM	323	CD	LYS	37	88.192	47.125	16.616	1.00 38.32	A_13
MOTA	324	CE	LYS	37	88.750	45.825	17.205	1.00 34.46	A_13
MOTA	325	NZ	LYS	37	88.234	44.576	16.557	1.00 12.49	A_13
ATOM	329	C	LYS	37	85.372	45.887	14.786	1.00 17.04	A_13
ATOM	330	0	LYS	37	86.131	44.958	15.053	1.00 18.14	A_13
MOTA	331	N	ALA	38	84.061	45.711	14.649	1.00 24.47	A_13
MOTA MOTA	333	CA	ALA	38	83.452	44.392	14.822	1.00 11.03	A_13
MOTA	334	CB	ALÀ	38	81.941	44.504	14.890	1.00 14.71	A_13
ATOM	335 336	Ç	ALA	38	83.900	43.451	13.697	1.00 20.27	A_13
MOTA		0	ALA	38	84.143	42.266	13.936	1.00 18.80	A_13
ATOM	339	N CA	PHE	39	84.021	43.971	12.477	1.00 22.58	A_13
MOTA	340	CB	PHE	39	84.492	43.158	11.355	1.00 18.87	A_13
ATOM	341	CG	PHE	39 39	84.350	43.899	10.027	1.00 19.91	A_13
MOTA	342		PHE	39	82.993	43.783	9.414	1.00 10.00	A_13
ATOM	343		PHE	39	82.266 82.438	44.915	9.097	1.00 17.54	A_13
ATOM	344		PHE	39	81.008	42.533 44.808	9.143 8.520	1.00 15.92	A_13
ATOM	345		PHE	39	81.186	42.418	8.569	1.00 20.75	A_13
ATOM	346	CZ	PHE	39	80.467	43.555	8.252	1.00 10.00	A_13
MOTA	347	C	PHE	39	85.955	42.827	11.589	1.00 10.00	A_13
ATOM	348	0	PHE	39	86.382	41.689	11.387	1.00 19.70	A_13 A_13
ATOM	349	N	LYS	40	86.699	43.822	12.072	1.00 21.31	A_13
ATOM	351	CA	LYS	40	88.117	43.673	12.369	1.00 20.07	A_13
MOTA	352	CB	LYS	40	88.703	44.967	12.927	1.00 13.77	A_13
ATOM	353	CG	LYS	40	90.192	44.885	13.171	1.00 11.54	A_13
MOTA	354	CD	LYS	40	90.757	46.242	13.507	1.00 10.34	A 13
MOTA	355	CE	LYS	40	92.236	46.142	13.838	1.00 11.24	A_13
ATOM	356	NZ	LYS	40	92.468	45.518	15.179	1.00 27.33	A_13
MOTA MOTA	360	C	LYS	40	88.352	42.534	13.337	1.00 12.06	A_13
ATOM	361 362	0	LYS	40	89.252	41.719	13.124	1.00 25.09	A_13
ATOM	364	N CA	VAL	41	87.495	42.418	14.349	1.00 12.26	A_13
MOTA	365	CB	VAL VAL	41	87.630	41.331	15.325	1.00 17.89	A_13
ATOM	366		VAL	41	86.351	41.205	16.216	1.00 10.00	A_13
ATOM	367		VAL	41 41	86.298	39.865	16.894	1.00 23.82	A_13 A_13
ATOM	368	C	VAL	41	86.329	42.274	17.259	1.00 17.65	A_13
ATOM	369	ŏ	VAL	41	87.822	40.009	14.560	1.00 23.06	A_13
ATOM	370	N	TRP	42	88.664	39.168	14.912	1.00 11.82	A_13
MOTA	372	CA	TRP	42	87.069 87.085	39.871	13.471	1.00 21.42	A_13
ATOM	373	CB	TRP	42	85.713	38.666 38:476	12.661	1.00 21.32	A_13
MOTA	374	CG	TRP	42	84.605	38.387	12.009	1.00 18.84	A_13
MOTA	375		TRP	42	84.437	37.369	13.025	1.00 25.92	A_13
MOTA	376		TRP	42	83.260	37.680	14.024 14.737	1.00 16.65	A_13
MOTA	377		TRP	42	85.165	36.223	14.737	1.00 17.58 1.00 11.14	A_13
MOTA	378		TRP	42	83.563	39.249	13.179	1.00 11.14	A_13 A_13
MOTA	379		TRP	42	82.755	38.832	14.200	1.00 10.00	A_13 A_13
MOTA	381	CZ2	TRP	42	82.785	36:879	15.793	1.00 10.91	A_13
MOTA	382	CZ3	TRP	42	84.691	35.425	15.436	1.00 23.68	A_13
MOTA	383	CH2	TRP	42	83.513	35.759	16.125	1.00 12.75	A_13
MOTA	384	C	TRP	42	88.190	38.600	11.623	1.00 27.45	A_13
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ATOM	385	0	TRP	42	88.834	37.556	11.472	1.00 11.84	A_13
ATOM	386		SER	43	88.413	39.702	10.909	1.00 25.46	A_13
ATOM	388	CA	SER	43	89.449	39.740	9.881	1.00 19.61	A_13
MOTA	389	CB	SER	43	89.342	40.993	8.991	1.00 16.16	
ATOM	390	OG	SER	43	89.495	42.199	9.709	1.00 26.34	A_13
MOTA	392				90.837	39.615			A_13
		Č	SER	43			10.491	1.00 11.53	A_13
ATOM	393	0	SER	43	91.758	39.119	9.834	1.00 17.99	A_13
MOTA	394	N	ASP	44	90.949	39.973	11.771	1.00 10.00	A_13
MOTA	396		ASP	44	92.206	39.908	12.505	1.00 16.90	A_13
MOTA	397	CB	ASP	44	92.057	40.588	13.857	1.00 17.79	A_13
ATOM	398	CG	ASP	44	92.544	42.013	13.839	1.00 15.93	A_13
MOTA	399	OD1	ASP	44	92.605	42.618	14.920	1.00 17.21	A_13
ATOM	400	OD2		44	92.874	42.533	12.754	1.00 19.50	A_13
ATOM	401	c	ASP	44	92.781	38.523	12.729	1.00 26.12	A_13
ATOM	402	Õ	ASP	44	93.996	38.362	12.897	1.00 21.21	A_13
ATOM	403	N.	VAL	45	91.911	37.523	12.745	1.00 20.89	A_13
ATOM	405	CA	VAL	45	92.353	36.161	12.996		A_13 A_13
	406	CB	VAL					1.00 27.53	
MOTA				45	91.853	35.678	14.381	1.00 16.30	A_13
MOTA	407	CG1		45	92.557	36.472	15.504	1.00 10.00	A_13
MOTA	408		VAL	. 45	90.348	35.857	14.495	1.00 10.86	A_13
ATOM	409	С	VAL	45	91.928	35.187	11.911	1.00 24.33	A_13
ATOM	410	0	VAL	45	91.864	33.978	12.157	1.00 18.84	A_13
MOTA	411	N	THR	46	91.750	35.705	10.694	1.00 16.30	A_13
MOTA	413	CA	THR	46	91.293	34.893	9.574	1.00 14.48	A_13
MOTA	414	CB	THR	46	89.750	34.796	9.662	1.00 22.05	A_13
MOTA	415	OG1	THR	46	89.279	33.609	9.028	1.00 31.53	A_13
MO:TA	417	CG2	THR	46	89.112	36.014	9.040	1.00 10.99	A_13
MOTA	418	С	THR	46	91.716	35.575	8.257	1.00 25.10	A_13
MOTA	419	0	THR	46	92.022	36.764	8.256	1.00 17.64	A_13
ATOM	420	N	PRO	47	91.688	34.845	7.114	1.00 15.31	A_13
ATOM	421	CD	PRO	47	91.459	33.398	6.985	1.00 17.94	A_13
MOTA	422	CA	PRO	47	92.069	35.416	5.815	1.00 21.50	A_13
MOTA	423	CB	PRO	47	92.199	34.182	4.911	1.00 21.50	
MOTA	424	CG	PRO	47	92.369				A_13
						33.041	5.848	1.00 27.45	A_13
MOTA	425	C	PRO	47	90.991	36.348	5.256	1.00 21.44	A_13
MOTA	426	0	PRO	47	91.095	36.788	4.116	1.00 11.08	A_13
MOTA	427	N	LEU	48	89.918	36.567	6.018	1.00 10.00	A_13
ATOM	429	CA	LEU	48	88.826	37.434	5.581	1.00 22.09	A_13
MOTA	430	CB	LEU	48	87.575	37.212	6.432	1.00 15.92	A_13
MOTA	431	CG	LEU	48	86.848	35.867	6.435	1.00 13.58	A_13
MOTA	432	CD1	LEU	48	85.931	35.811	7.654	1.00 25.90	A_13
MOTA	433	CD2	LEU	48	86.073	35.666	5.157	1.00 16.47	A_13
MOTA	434	С	LEU	48	89.156	38.916	5.641	1.00 21.20	A_13
MOTA	435	0	LEU	48	89.936	39.366	6.480	1.00 17.28	A_13
ATOM	436	N	ASN	49	88.569	39.670	4.723	1.00 26.12	A_13
ATOM	438	CA	ASN	49	88.738	41.112	4.717	1.00 26.84	A_13
ATOM	439	CB	ASN	49	89.936	41.569	3.885	1.00 18.29	A_13
ATOM	440	CG	ASN	49	90.010	40.912	2.568	1.00 22.55	A_13
ATOM	441		ASN	49	90.928	40.131	2.305		A_13
ATOM	442		ASN	49				1.00 24.41	A_13
ATOM	445	C		49	89.068	41.235	1.693	1.00 46.51	A_13
ATOM			ASN		87.416	41.705	4.259	1.00 12.18	A_13
	446	0	ASN	49	86.732	41.128	3.400	1.00 20.77	A_13
ATOM	447	N	PHE	50	87.025	42.802	4.900	1.00 21.39	A_13
MOTA	449	CA	PHE	50	85.738	43.439	4.642	1.00 10.00	A_13
MOTA	450	CB	PHE	50	84.914	43.440	5.932	1.00 11.45	A_13
ATOM	451	CG	PHE	50	84.863	42.098	6.629	1.00 10.63	A_13
MOTA	452		PHE	50	85.886	41.705	7.490	1.00 10.00	A_13
MOTA	453		PHE	50	83.809	41.216	6.395	1.00 14.63	A_13
MOTA	454		PHE	50	85.858	40.457	8.097	1.00 26.88	A_13
MOTA	455	CE2	PHE	50	83.773	39.963	7.000	1.00 21.13	A 13
MOTA	456	CZ	PHE	50	84.801	39.581	7.852	1.00 10.30	A 13
MOTA	457	С	PHE	50	85.867	44.842	4.093	1.00 22.56	A_13 A_13
ATOM	458	0	PHE	50	86.638	45.644	4.612	1.00 19.33	A_13
ATOM	459	N	THR	51	85.099	45.129	3.044	1.00 21.47	A_13
ATOM	461	CA	THR	51	85.125	46.433	2.371	1.00 24.21	7 13
MOTA	462	CB	THR	51	85.602				A_13
ATOM	463					46.306	0.895	1.00 15.39	A_13
			THR	51	86.950	45.811	0.853	1.00 24.33	A_13
MOTA	465		THR	51	85.551	47.654	0.192	1.00 25.47	A_13
MOTA	466	C	THR	51	83.735	47.048	2.359	1.00 22.17	A_13
ATOM	467	0	THR	51	82.766	46.421	1.912	1.00 20.53	A_13
ATOM	468	N	ARG	52	83.653	48.294	2.797	1.00 16.53	A_13
MOTA	470	CA	ARG	52	82.393	49.004	2.871	1.00 10.00	A_13
MOTA	471	CB	ARG	52	82.490	50.085	3.939	1.00 10.00	A_13
MOTA	472	CG	ARG	52	81.201	50.778	4.259	1.00 12.47	A_13
ATOM	473	CD	ARG	52	81.462	51.879	5.278	1.00 19.61	A_13
MOTA	474	NE	ARG	52	80.371	52.836	5.333	1.00 30.55	A_13
MOTA	476	CZ	ARG	52	80.489		5.795	1.00 24.06	A_13
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ATOM	477	NH1	ARC	52	81.661	54.508	6.257	1.00 21.24	2 12
									A_13
MOTA	480	NH2	ARG	52	79.421	54.862	5.829	1.00 27.78	A_13
MOTA	483	С	ARG	52	81.980	49.620	1.540	1.00 30.22	A_13
ATOM	484	0	ARG	52	82.782	50.269	0.859	1.00 16.27	A_13
MOTA	485	N	LEU	53	80.730	49.372	1.161	1.00 21.07	A_13
ATOM	487	CA	LEU	53	80.159	49.914	-0.062	1.00 15.73	A_13
ATOM	488	CB	LEU	53	79.435	48.831	-0.868	1.00 11.53	A_13
ATOM	489	CG	LEU						
				53	80.304	47.770	-1.530	1.00 10.00	A_13
MOTA	490	CD1	LEU	53	79:429	46.790	-2.296	1.00 13.21	A_13
MOTA	491	CD2	LEU	53	81.280	48.443	-2.448	1.00 12.78	A_13
ATOM	492	C	LEU	53	79.149	50.932	0.421		7-13
								1.00 10.00	A_13
MOTA	493	0	LEU	53	78.463	50.713	1.411	1.00 13.62	A_13
MOTA	494	N	HIS	54	79.043	52.041	-0.283	1.00 15.73	A_13
ATOM	496	CA	HIS .		78.102	53.065	0.126	1.00 12.47	
									A_13
MOTA	497	CB	HIS	54	78.765	54.435	0.011	1.00 15.18	A_13
MOTA	498	CG	HIS	54	79.967	54.589	0.884	1.00 21.27	A_13
MOTA	499	CD2	HIS	54	81.207	54.056	0.798	1.00 25.30	A_13
ATOM	500		HIS	54	79.951	55.338			
							2.043	1.00 16.48	A_13
MOTA	502		HIS	54	81.127	55.255	2.633	1.00 21.62	A_13
ATOM	503	NE2	HIS	54	81.910	54.482	1.899	1.00 29.91	A_13
ATOM	505	С	HIS	54	76.796	53.044	-0.664	1.00 15.50	A_13
ATOM	506	ŏ	HIS	54	75.914				2-13
						53.849	-0.403	1.00 21.80	A_13
ATOM	507	N	ASP	55	76.707	52.178	-1.671	1.00 18.31	A_13
ATOM	509	CA	ASP	55	75.509	52.077	-2.502	1.00 17.23	A_13
ATOM	510	CB	ASP	55	75.645	52.928	-3.773	1.00 19.94	
									A_13
MOTA	511	CG	ASP	55	75.864	54.393	-3.495	1.00 26.81	A_13
MOTA	512	OD1	ASP	55	75.059	54.991	-2.741	1.00 35.97	A_13
ATOM	513	002	ASP	55	76.839	54.948	-4.058	1.00 25.09	A_13
	514								7-13
MOTA		C	ASP	55	75.343	50.645	-2.970	1.00 21.50	A_13
MOTA	515	0	ASP	55	76.286	49.862	-2.929	1.00 17.45	A_13
ATOM	516	N	GLY	56	74.160	50.337	-3.489	1.00 10.31	A_13
ATOM	518	CA	GLY	56	73.897	49.014	-4.014	1.00 13.67	
									A_13
MOTA	519	С	GLY	56	73.842	47.869	-3.030	1.00 17.61	A_13
ATOM	520	0	GLY	56	73.683	48.065	-1.825	1.00 12.57	A_13
MOTA	521	N	ILE	57	73.943	46.653	-3.560	1.00 22.27	A_13
									V-13
ATOM .	523	CA	ILE	57	73.895	45.460	-2.737	1.00 11.39	A_13
ATOM	524	CB	ILE	57	72.941	44.391	-3.347	1.00 22.87	A_13
ATOM	525	CG2	ILE	57	73.365	42.995	-2.955	1.00 22.98	A_13
ATOM	526								
			ILE	. 57	71.522	44.582	-2.787	1.00 30.87	A_13
MOTA	527	CD1	ILE	57	71.002	46.022	-2.796	1.00 28.15	A_13
MOTA	528	С	ILE	57	75.289	44.919	-2.446	1.00 22.32	A_13
ATOM	529	Ō	ILE	57	76.140	44.849	-3.332		
					70.140			1.00 25.00	A_13
MOTA	530	N	ALA	58	75.517	44.631	-1.168	1.00 25.02	A_13
MOTA	532	CA	ALA	58	76.773	44.105	-0.669	1.00 15.45	A_13
MOTA	533	CB	ALA	58	77.366	45.060	0.358	1.00 11.62	A_13
MOTA	534	C	ALA	58	76.438	42.780	-0.006	1.00 12.08	A_13
MOTA	535	0	ALA	58	75.289	42.521	0.307	1.00 13.30	A_13
ATOM	536	N	ASP	59	77.449	41.968	0.247	1.00 14.79	A_13
ATOM	538	CA	ASP	59	77.245	40.675	0.880		7 1 2
									A_13
ATOM	539	CB	ASP	59	78.608	39.974	1.093	1.00 10.83	A_13
ATOM	540	CG	ASP	59	79.425	39.858	-0.210	1.00 23.35	A_13
ATOM	541	OD1	ASP	59	80.598	40.266	-0.236	1.00 17.98	A_13
MOTA	542			59	78.896				
			ASP			39.379	-1.230	1.00 16.89	A_13
MOTA	543	C	ASP	59	76.480	40.806	2.200	1.00 13.69	A_13
MOTA	544	0	ASP	59	75.402	40.227	2.380	1.00 15.93	A_13
ATOM	545	N	ILE	60	77.025	41.596	3.109	1.00 13.15	A_13
MOTA	547	CA	ILE	60	76.422				
						41.800	4.412	1.00 12.20	A_13
MOTA	548	CB	ILE	60	77.500	41.695	5.508	1.00 12.12	A_13
ATOM	549	CG2	ILE	60	76.921	42.060	6.864	1.00 19.27	A_13
MOTA	550	CG1	ILE	60	78.118	40.287	5.481	1.00 10.00	A_13
ATOM	551		ILE						A_13
				60	79.330	40.120	6.360	1.00 10.00	A_13
MOTA	552	С	ILE	60	75.743	43.164	4.456	1.00 17.78	` A_13
ATOM	553	0	ILE	60	76.410	44.193	4.478	1.00 18.65	A_13
ATOM	554	N	MET	61	74.416				2-13
						43.168	4.431	1.00 12.54	A_13
ATOM	556	CA	MET	61	73.640	44.416	4.476	1.00 12.86	A_13
MOTA	557	CB	MET	61	72.385	44.314	3.604	1.00 18.16	A_13
MOTA	558	. CG	MET	61	72.634	43.979			2 13
							2.141	1.00 10.00	A_13
ATOM	559	SD	MET	61	73.374	45.314	1.251	1.00 10.69	A_13
ATOM	560	CE	MET	61	71.836	46.299	0.764	1.00 10.00	A_13
ATOM	561	C	MET	61	73.239	44.666	5.921		A_13
								1.00 10.15	
ATOM	562	0	MET	61	72.584	43.838	6.547	1.00 18.13	A_13
MOTA	563	N	ILE	62	73.706	45.784	6.456	1.00 15.60	A_13
MOTA	565	CA	ILE	62	73.452	46.170	7.837	1.00 18.55	A_13
ATOM	566		ILE						W-13
		CB		62	74.723	46.828	8.437	1.00 10.00	A_13
MOTA	567		ILE	62	74.498	47.163	9.900	1.00 26.36	A_13
MOTA	568	CG1	ILE	62	75.936	45.897	8.302	1.00 11.04	A_13
MOTA	569		ILE	62	77.228	46.481	8.891	1.00 10.00	A_13
						-0.401	0.071	2.00 10.00	Y_73

ATOM	570	С	ILE	62	72.289	47.172	7.920	1.00 17.99	A_13
ATOM	571		ILE	62	72.335	48.208	7.264	1.00 12.72	A_13
ATOM ATOM	572 574		SER SER	63. 63	71.285 70.149	46.896 47.803	8.751 8.882	1.00 10.00 1.00 12.52	A_13
ATOM	575		SER	63	69.016	47.364	7.956	1.00 12.52	A_13 A_13
ATOM	576		SER	63	68.448	46.146	8.415	1.00 27.90	A_13
MOTA	578		SER	63	69.625	47.854	10.314	1.00 13.14	A_13
ATOM	579	-	SER	63	69.869	46.951	11.101	1.00 22.10	A_13
ATOM ATOM	580	N	PHE PHE	64 64	68.919	48.932	10.640	1.00 21.17	A_13
ATOM	582 583	CA CB	PHE	64	68.317 68.777	49.139 50.468	11.954 12.574	1.00 22.01 1.00 10.98	A_13 A_13
ATOM	584	CG	PHE	64	70.189	50.448	13.092	1.00 10.00	A_13
ATOM	585	CD1		64	70.473	49.885	14.322	1.00 10.00	A_13
MOTA	586	CD2		64	71.229	51.016	12.357	1.00 16.56	A_13
ATOM	587	CE1		64	71.777	49.885	14.825	1.00 10.00	A_13
MOTA MOTA	588 589	CE2 CZ	PHE PHE	64 64	72.540 72.812	51.025 50.459	12.846 14.081	1.00 10.00 1.00 18.83	A_13 A_13
MOTA	590	C	PHE	64	66.825	49.207	11.675	1.00 22.55	A_13 A_13
MOTA	591	0	PHE	64	66.405	49.940	10.779	1.00 19.49	A_13
ATOM	592	N	GLY	65	66.031	48.485	12.453	1.00 13.69	A_13
MOTA	594	CA	GLY	65 65	64.593	48.491	12.238	1.00 10.70	A_13
MOTA MOTA	595 596	C O	GLY GLY	65 65	63.894 64.559	48.138 47.777	13.521 14.491	1.00 12.62 1.00 18.29	A_13 A_13
ATOM	597	N	ILE	66	62.577	48.309	13.565	1.00 13.69	A_13
MOTA	599	CA	ILE	66	61.803	47.968	14.760	1.00 21.58	A_13
MOTA	600	CB	ILE	66	61.227	49.228	15.503	1.00 30.51	A_13
ATOM ATOM	601 602	CG2 CG1		66 66	62.351	50.110	16.025	1.00 10.43	A_13
ATOM	603	CD1	ILE	66 66	60.332 59.587	50.062 51.149	14.586 15.333	1.00 14.56 1.00 16.94	A_13 A_13
ATOM	604	C	ILE	66	60.662	47.030	14.361	1.00 10.94	A_13
ATOM	605	0	ILE	66	60.311	46.962	13.188	1.00 10.00	A_13
ATOM	606	N	LYS	67	60.143	46.271	15.330	1.00 10.00	A_13
ATOM	608	CA	LYS	67 67	59.036	45.327	15.103	1.00 10.23	A_13
MOTA MOTA	609 610	CB	LYS	67 67	57.689 57.584	46.042 46.895	15.268 16.510	1.00 10.29 1.00 14.63	A_13 A_13
ATOM	611	CD	LYS	67	57.646	46.056	17.774	1.00 14.03	A_13
MOTA	612	CE	LYS	67	57.382	46.923	18.986	1.00 22.99	A_13
MOTA	613	NZ	LYS	67	57.480	46.174	20.258	1.00 28.27	A_13
MOTA	617 618	C	LYS	67 67	59.113	44.633	13.726	1.00 17.91	A_13
MOTA MOTA	619	N O	LYS GLU	67 68	60.167 58.027	44.106 44.690	13.366 12.949	1.00 24.16 1.00 12.72	A_13 A_13
ATOM	621	CA	GLU	68	57.960	44.067	11.624	1.00 16.06	A_13 A_13
MOTA	622	CB	GLU	68	56.505	44.019	11.128	1.00 26.89	A_13
ATOM	623	CG	GLU	68	55.566	43.258	12.087	1.00 36.97	A_13
MOTA MOTA	624 625	CD OF1	GLU GLU	68 68	54.217	43.973	12.381	1.00 41.61	A_13
ATOM	626		GLU	68 68	53.289 54.074	43.921 44.561	11.537 13.485	1.00 17.31 1.00 26.72	A_13 A_13
ATOM	627	c	GLU	68	58.823	44.911	10.705	1.00 22.50	A_13
ATOM	628	0	GLU	68	58.587	46.093	10.532	1.00 20.64	A_13
ATOM	629	N	HIS	69 60	59.848	44.315	10.120	1.00 16.43	A_13
MOTA MOTA	631 632	CA CB	HIS HIS	69 69	60.732 61.930	45.102 45.603	9.283 10.103	1.00 13.69 1.00 10.97	A_13 A_13
ATOM	633	CG	HIS	69	62.786	44.502	10.643	1.00 24.02	A_13 A_13
ATOM	634		HIS	69	63.873	43.876	10.133	1.00 10.00	A_13
ATOM	635		HIS	69	62.512	43.876	11.839	1.00 17.68	A_13
ATOM ATOM	637 638		HIS HIS	69 69	63.384 64.228	42.912	12.041 11.020	1.00 12.53	A_13
ATOM	639	C	HIS	69	61.214	42.888 44.469	7.983	1.00 10.00 1.00 21.28	A_13 A_13
ATOM	640	0	HIS	69	62.314	44.780	7.529	1.00 18.74	A_13
MOTA	641	N	GLY	70	60.451	43.537	7.411	1.00 13.11	A_13
MOTA MOTA	643 644	CA	GLY	70	60.832	42.968	6.127	1.00 10.00	A_13
ATOM	645	CO	GLY GLY	70 70	61.262 61.523	41.533 41.125	5.936 4.794	1.00 10.00 1.00 15.12	A_13
MOTA	646	N	ASP	71	61.412	40.768	7.012	1.00 19.99	A_13 A_13
MOTA	648	CA	ASP	71	61.842	39.381	6.862	1.00 19.99	A_13
MOTA	649	CB	ASP	71	63.332	39.223	7.218	1.00 10.00	A_13
MOTA	650	CG	ASP	71	63.672	39.752	8.592	1.00 23.52	A_13
MOTA MOTA	651 652		ASP	71 71	64.846	40.110	8.803	1.00 13.38	A_13
ATOM	653	C	ASP	71	62.774 60.998	39.812 38.377	9.464 7.632	1.00 12.94 1.00 22.07	A_13 A_13
ATOM	654	ŏ	ASP	71	61.319	37.190	7.649	1.00 24.45	A_13
MOTA	655	N	PHE	72	59.946	38.865	8.292	1.00 14.15	A_13
ATOM	657	CA	PHE	72	59.040	38.035	9.094	1.00 10.00	A_13
MOTA MOTA	658 659	CB CG	PHE	72 72	58.410	36.905	8.272	1.00 10.00	A_13
ATOM	660		PHE	72	57.360 56.115	37.387 37.773	7.332 7.815	1.00 10.00 1.00 23.01	A_13 A_13
MOTA	661		PHE	72	57.624	37.507	5.973	1.00 12.52	A_13
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ATOM	662	CE1	PHE	72	55.144	38.290	6.950	1.00 18.99	A_13
MOTA	663		PHE	72	56.662	38.023	5.091	1.00 13.37	A_13
MOTA	664	CZ	PHE	72	55.420	38.413	5.576	1.00 22.50	A_13
MOTA	665	С	PHE	72	59.634	37.523	10.392	1.00 16.31	A_13
ATOM	666	0	PHE	72	59.111	36.596	11.021	1.00 15.64	A_13
MOTA	667	N	TYR	73	60.737	38.141	10.793	1.00 18.10	A_13
	669		TYR	73	61.407	37.827		1.00 14.01	A_13
MOTA							12.046		A_13
MOTA	670	CB	TYR	73	62.845	37.331	11.803	1.00 21.08	A_13
ATOM	671	CG	TYR	73	62.915	35.965	11.138	1.00 22.48	A_13
MOTA	672	CD1	TYR	73	63.579	35.788	9.923	1.00 30.23	A_13
			TYR	73	63.615		9.291		
ATOM	.673					34.538		1.00 24.04	A_13
MOTA	674		TYR	73	62.288	34.856	11.710	1.00 19.23	A_13
MOTA	675	CE2	TYR	73	62.320	33.606	11.083	1.00 29.35	A_13
ATOM	676	CZ	TYR	73	62.984	33.460	9.875	1.00 12.50	A_13
ATOM	677	OH .	TYR	73	63.018	32.246	9.241	1.00 17.89	A_13
									N_13
ATOM	679	С	TYR	73	61.360	39.203	12.721	1.00 22.00	A_13
ATOM	680	0	TYR	73	62.365	39.919	12.819	1.00 10.93	A_13
MOTA	681	N	PRO	74	60.175	39.570	13.221	1.00 19.94	A_13
ATOM	682	CD	PRO	74	58.969	38.723	13.278	1.00 15.69	A_13
	683								
MOTA		CA	PRO	74	59.934	40.843	13.886	1.00 16.75	A_13
ATOM	684	CB	PRO	74	58.417	40.836	14.067	1.00 17.27	A_13
ATOM	685	CG	PRO	74	58.131	39.407	14.335	1.00 16.24	A_13
ATOM	686	С	PRO	74	60.640	41.037	15.216	1.00 17.39	A_13
ATOM	687	ō	PRO	74	60.779	40.105	16.023	1.00 10.00	A_13
									A_13
ATOM	688	N	PHE	75	61.098	42.264	15.431	1.00 10.00	A_13
MOTA	690	CA	PHE	75	61.743	42.618	16.675	1.00 16.45	A_13
ATOM	691	CB	PHE	75	62.613	43.865	16.512	1.00 20.71	A_13
MOTA	692	CG	PHE	75	63.931	43.590	15.841	1.00 23.32	A_13
	693		PHE		64.694				
MOTA				75		42.482	16.200	1.00 12.03	A_13
MOTA	694		PHE	75	64.405	44.420	14.842	1.00 22.30	A_13
ATOM	695	CE1	PHE	75	65.905	42.214	15.572	1.00 17.64	A_13
MOTA	696	CE2	PHE	75	65.622	44.148	14.208	1.00 15.43	A_13
ATOM	697	cz	PHE	75	66.367	43.044	14,576	1.00 10.00	A_13
MOTA	698	C	PHE	75	60.632	42.784	17.707	1.00 25.73	A_13
ATOM	699	0	PHE	75	59.443	42.778	17.370	1.00 18.57	A_13
MOTA	700	N	ASP	76	61.009	43.002	18.952	1.00 20.50	A_13
ATOM	702	CA	ASP	76	60.023	43.049	20.006	1.00 13.89	A_13
	703		ASP						
MOTA		CB		76	60.241	41.805	20.873	1.00 20.69	A_13
ATOM	704	CG	ASP	76	61.672	41.685	21.378	1.00 22.52	A_13
ATOM	705	OD1	ASP	76	61.947	40.771	22.174	1.00 20.06	A_13
MOTA	706	OD2	ASP	76	62.525	42.506	20.998	1.00 10.69	A_13
ATOM	707	c	ASP	76	59.971	44.277			7.13
							20.900	1.00 25.20	A_13
MOTA	708	0	ASP	76	59.397	44.207	21.986	1.00 29.52	A_13
MOTA	709	N	GLY	77	60.585	45.379	20.488	1.00 10.00	A_13
MOTA	711	CA	GLY	77	60.575	46.553	21.334	1.00 10.00	A_13
ATOM	712	C	GLY	77	61.769	46.514	22.266	1.00 10.00	A_13
ATOM	713	ŏ	GLY	77					M_13
					62.735	45.797	21.987	1.00 18.49	A_13
MOTA	714	N	PRO	78	61.785	47.344	23.322	1.00 16.07	A_13
MOTA	715	CD	PRO	78	60.790	48.426	23.505	1.00 15.88	A_13
MOTA	716	CA	PRO	78	62.855	47.439	24.330	1.00 16.23	A_13
MOTA	717	CB	PRO	78	62.261	48.391	25.363	1.00 22.96	A_13
ATOM	718	CG	PRO	78	61.470	49.349			2-13
							24.501	1.00 22.37	A_13
MOTA	719	C	PRO	78	63.150	46.090	24.969	1.00 25.32	A_13
MOTA	720	0	PRO	78	62.227	45.356	25.272	1.00 20.04	A_13
MOTA	721	N	SER	79	64.432	45.750	25.099	1.00 20.93	A_13
ATOM	723	CA	SER	79	64.878	44.478	25.689	1.00 20.51	A_13
ATOM	724	CB	SER	79	64.364	44.311	27.131	1.00 23.69	
MOTA	725	OG							A_13
			SER	79	65.028	45.211	28.006	1.00 33.37	A_13
MOTA	727	С	SER	79	64.557	43.248	24.863	1.00 20.39	A_13
MOTA	728	0	SER	79	64.124	43.362	23.708	1.00 17.27	A_13
MOTA	729	N	GLY	80	64.825	42.071	25.415	1.00 13.38	A_13
ATOM	731	CA	GLY	80	64.564	40.850	24.678		
								1.00 10.11	A_13
MOTA	732	C	GLY	80	65.471	40.808	23.458	1.00 13.15	A_13
MOTA	733	0	GLY	80	66.614	41.251	23.538	1.00 31.80	A_13
MOTA	734	N	LEU	81	64.939	40.393	22.310	1.00 29.05	A_13
ATOM	736	CA	LEU	81	65.720	40.317	21.078	1.00 29.63	A_13
MOTA	737	CB	LEU	81					W_13
					64.789	40.033	19.905	1.00 19.67	A_13
MOTA	738	CG	LEU	81	65.121	38.872	18.971	1.00 21.79	A_13
MOTA	739		LEU	81	64.215	38.980	17.773	1.00 23.87	A_13
ATOM	740		LEU	81	66.590	38.918	18.518	1.00 22.09	A_13
MOTA	741	C	LEU	81	66.442	41.649			
	742						20.835	1.00 19.25	A_13
MOTA		0	LEU	81	65.808	42.700	20.872	1.00 14.95	A_13
ATOM	743	N	LEU	82	67.760	41.599	20.657	1.00 25.03	A_13
ATOM	745	CA	LEU	82	68.573	42.795	20.421	1.00 27.35	A_13
ATOM	746	CB	LEU	82	69.868	42.747	21.244	1.00 12.74	A_13
ATOM	747	CG	LEU		69.802	42.748			
ATOM	748		LEU				22.773	1.00 16.50	A_13
ALUM	740	CDI	PEU	02	68.590	43.520	23.263	1.00 17.99	A_13

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ATOM	749	CD2		82	69.744	41.343	23.279	1.00 13.28	A_13
ATOM	750	Č	LEU	82	68.938	42.945	18.949	1.00 24.79	A_13
MOTA	751	0	LEU	82	68.812	44.039	18.363	1.00 14.36	A_13
ATOM	752	N	ALA	83	69.387	41.839	18.359	1.00 21.15	A_13
MOTA ATOM	754 755	CA	ALA	83 83	69.790 71.180	41.819	16.961	1.00 15.64	A_13
ATOM	756	CB C	ALA ALA	83	69.806	42.410 40.400	16.820	1.00 15.74 1.00 19.37	A_13
ATOM	757	ŏ	ALA	83	69.864	39.458	16.444 17.227	1.00 19.37	A_13
ATOM	758	N	HIS	84	69.746	40.252	15.126	1.00 20.42	A_13 A_13
ATOM	760	CA	HIS	84	69.808	38.939	14.502	1.00 20.51	A_13 A_13
ATOM	761	CB	HIS	84	68.454	38.185	14.476	1.00 12.34	A_13
ATOM	762	CG	HIS	84	67.361	38.849	13.679	1.00 24.79	A_13
ATOM	763	CD2		84	67.381	39.489	12.488	1.00 10.00	A_13
MOTA	764	ND1		84	66.052	38.869	14.104	1.00 13.50	A_13
MOTA	766	CE1		84	65.307	39.497	13.210	1.00 14.37	A_13
ATOM	767	NE2		84	. 66.087	39.886	12.220	1.00 15.00	A_13
MOTA	768	С	HIS	84	70.418	39.088	13.130	1.00 22.78	A_13
ATOM	769	0	HIS	84	70.338	40.162	12.532	1.00 10.00	A_13
ATOM	770	N	ALA	85	71.086	38.027	12.685	1.00 13.43	A_13
ATOM	772	CA	ALA	85	71.746	37.983	11.402	1.00 10.00	A_13
ATOM	773	CB	ALA	85	73.234	38.132	11.596	1.00 10.05	A_13
ATOM	774	Ç	ALA	85	71.426	36.661	10.721	1.00 17.89	A_13
MOTA	775 776	0	ALA	85 86	70.900	35.746	11.346	1.00 19.43	A_13
ATOM ATOM	778	N CA	PHE	86 86	71.697	36.585	9.425 8.651	1.00 13.49	À_13
ATOM	779	CB	PHE	86	71.459 70.739	35.372 35.728	7.344	1.00 12.49	A_13
ATOM	780	CG	PHE	86	69.348	36.240	7.529	1.00 10.00 1.00 19.96	A_13 A_13
ATOM	781		PHE	86	68.252	35.434	7.212	1.00 21.89	A_13
MOTA	782		PHE	86	69.119	37.530	8.003	1.00 10.63	A_13
ATOM	783	CE1		86	66.946	35.900	7.364	1.00 16.59	A_13
ATOM	784	CE2	PHE	86	67.829	38.009	8.158	1.00 19.06	A_13
MOTA	785	CZ	PHE	86	66.732	37.194	7.838	1.00 24.79	A_13
ATOM	786	C	PHE	86	72.802	34.721	8.298	1.00 11.05	A_13
ATOM	787	0	PHE	86	73.774	35.435	8.041	1.00 25.56	A_13
MOTA	788	N	PRO	87	72.892	33.375	8.304	1.00 19.41	A_13
ATOM	789	CD	PRO	87	71.876	32.383	8.717	1.00 17.25	A_13
ATOM	790	CA	PRO	87	74.149	32.686	7.956	1.00 29.29	A_13
ATOM	791	CB	PRO	87	73.800	31.198	8.135	1.00 18.88	A_13
MOTA	792 793	CG	PRO	87 87	72.329	31.160	7.939	1.00 20.17	A_13
MOTA MOTA	794	C	PRO	87 87	74.562 73.728	32.999 33.448	6.503	1.00 10.00	A_13
ATOM	795	N	PRO	88	75.814	32.701	5.703 6.120	1.00 20.68	A_13 A_13
ATOM	796	CD	PRO	88	76.796	31.854	6.831	1.00 10.00 1.00 19.58	A_13 A_13
ATOM	797	CA	PRO	88	76.280	32.977	4.756	1.00 12.43	A_13
MOTA	798	СВ	PRO	88	77.600	32.201	4.676	1.00 18.69	A_13
ATOM	799	CG	PRO	88	78.073	32.163	6.098	1.00 18.48	A_13
MOTA	800	C	PRO	88	75.304	32.510	3.672	1.00 24.39	A_13
MOTA	801	0	PRO	88	74.596	31.522	3.854	1.00 16.92	A_13
ATOM	802	N	GLY	89	75.266	33.230	2.560	1.00 10.73	A_13
ATOM	804	CA	GLY	89	74.386	32.868	1.471	1.00 10.00	A_13
MOTA	805	C	GLY	89	73.960	34.127	0.772	1.00 10.94	A_13
MOTA	806	0	GLY	89	74.143	35.218	1.307	1.00 19.86	A_13
MOTA	807 808	N	PRO	90 90	73.390	34.019	-0.432	1.00 26.31	A_13
ATOM ATOM	809	CD CA	PRO PRO	90	73.090 72.960	32.792 35.212	-1.192 -1.163	1.00 18.46 1.00 25.07	A_13
ATOM	8,10	CB	PRO	90	72.670	34.651	-2.556		A_13
MOTA	811	CG	PRO	90	72.108	33.289	-2.236	1.00 15.47 1.00 24.63	A_13 A_13
MOTA	812	C	PRO	90	71.726	35.879	-0.543	1.00 20.41	A_13
ATOM	813	0	PRO	90	71.176	35.390	0.442	1.00 17.00	A_13
ATOM	814	N	ASN	91	71.303	37.000	-1.125	1.00 18.43	A_13
MOTA	816	CA	ASN	91	70.127	37.721	-0.653	1.00 14.03	A_13
MOTA	817	CB	ASN	91	68.863	36.932	-0.999	1.00 15.26	A_13
MOTA	818	CG	ASN	91	68.860	36.430	-2.439	1.00 36.74	A_13
MOTA	819		ASN	91	68.497	35.282	-2.701	1.00 29.56	A_13
ATOM	820		ASN	91	69.265	37.286	-3.376	1.00 27.03	A_13
MOTA	823	C	ASN	91	70.226	37.986	0.849	1.00 24.66	A_13
MOTA	824	0	ASN	91	71.257	38.479	1.313	1.00 17.43	A_13
MOTA MOTA	825 827	N CA	TYR	92 92	69.198	37.632	1.622	1.00 17.69	A_13
ATOM	828	CB	TYR. TYR	92 92	69.233 67.942	37.876	3.061	1.00 10.17	A_13
ATOM	829	CG	TYR	92	66.786	37.428 38.364	3.744 3.523	1.00 16.78 1.00 26.17	A_13
ATOM	830	CD1		92	66.015	38.803	4.581	1.00 26.17	A_13 A_13
MOTA	831	CEI		92	64.947	39.678	4.380	1.00 29.60	A_13
MOTA	832	CD2		92	66.467	38.818	2.250	1.00 25.90	A_13
MOTA	833	CE2	TYR	92	65.406	39.691	2.040	1.00 30.60	A_13
ATOM	834	CZ	TYR	92	64.647	40.117	3.107	1.00 12.31	A_13
MOTA	835	ОН	TYR	92	63.575	40.967	2.886	1.00 26.07	A_13
									•

ATOM	837	С	TYR	92	70.427	37.245	3.763	1.00 11.94	n 12
									A_13
MOTA	838		TYR	92	70.752	37.617	4.882	1.00 17.58	A_13
ATOM	839	N	GLY	93	71.095	36.311	3.097	1.00 24.67	A_13
ATOM	841		GLY	93	72.250	35.666	3.691	1.00 18.05	3-13
									A_13
ATOM	842	С	GLY	93	73.295	36.681	4.116	1.00 10.00	A_13
MOTA	843	0	GLY	93	73.573	37.656	3.391	1.00 10.13	A_13
ATOM	844	N	GLY	94	73.812	36.495	5.328	1.00 12.44	A_13
MOTA	846	CA	GLY	94	74.827	37.372	5.872	1.00 10.00	A_13
ATOM	847	С	GLY	94	74.358	38.694	6.456	1.00 17.29	A_13
ATOM	848	0	GLY	94	75.052	39.271	7.284	1.00 14.53	A_13
MOTA	849	N	ASP	95	73.221	39.206	5.993	1.00 10.00	A_13
MOTA	851	CA	ASP	95	72.689	40.485	6.472	1.00 16.35	A_13
MOTA	852	CB	ASP	95	71.332	40.777	5.814	1.00 10.00	A_13
MOTA	853	CG	ASP	95	71.421	40.904	4.309	1.00 14.54	A_13
ATOM	854	OD1		95	70.406	41.256	3.673	1.00 11.86	·
									A_13
ATOM	855	OD2	ASP	95	72.502	40.647	3.753	1.00 15.39	A_13
ATOM	856	С	ASP	95	72.548	40.523	7.994	1.00 22.31	A_13
						30.323			
MOTA	857	0	ASP	95	72.279	39.497	8.635	1.00 10.88	A_13
ATOM	858	N	ALA	96	72.703	41.711	8.566	1.00 18.45	A_13
ATOM	860	CA	ALA	96					7_13
					72.609	41.877	10.011	1.00 15.08	A_13
ATOM	861	CB	ALA	96	73.982	42.244	10.587	1.00 19.20	A_13
ATOM	862	C	ALA	96	71.587	42.961	10.345	1.00 14.91	A_13
MOTA	863	0	ALA	96	71.702	44.092	9.876	1.00 10.00	A_13
ATOM	864	N	HIS	97	70.635	42.646	11.215	1.00 14.01	À_13
									2-13
MOTA	866	CA	HIS	97	69.599	43.620	11.581	1.00 11.35	A_13
ATOM	867	CB	HIS	97	68.207	43.083	11.203	1.00 20.32	A_13
ATOM	868	CG	HIS	97	68.027	42.786			
							9.742	1.00 15.00	A_13
ATOM	869	CD2	HIS	97	68.734	43.186	8.654	1.00 10.00	A_13
MOTA	870	ND1	HTC	97	67.014	41.978	9.257	1.00 14.03	
									A_13
MOTA	871	CEI	HIS	97	67.108	41.895	7.936	1.00 10.00	A_13
ATOM	872	NE2	HTS	97	68.142	42.618	7.552	1.00 17.10	A_13
MOTA	874	C	HIS	97	69.650	43.952	13.078	1.00 13.37	A_13
MOTA	875	0	HIS	97	69.736	. 43.055	13.908	1.00 13.48	A_13
ATOM	876						12 422		
		И	PHE	98	69.596	45.237	13.423	1.00 21.01	A_13
ATOM	878	CA	PHE	98	69.634	45.668	14.823	1.00 11.27	A_13
ATOM	879	CB	PHE	98	70.817	46.615	15.055		
								1.00 10.00	A_13
MOTA	880	CG	PHE	98	72.138	46.011	14.703	1.00 20.49	A_13
ATOM	881	CD1	PHE	98	72.984	45.524	15.707	1.00 17.49	
				4 .					A_13
ATOM	882	CD2	PHE	98	72.506	45.853	13.365	1.00 13.51	A_13
ATOM	883	CEl	PHE	98	74.171	44.888	15.382	1.00 20.00	A_13
ATOM	884	CE2	PHE	98	73.693	45.215	13.024	1.00 10.00	A_13
ATOM	885	CZ	PHE	98	74.527	44.728	14.029	1.00 10.00	A_13
ATOM	886	c	PHE						
				98	68.336	46.336	15.245	1.00 25.38	A_13
ATOM	887	0	PHE	98	67.815	47.218	14.552	1.00 10.00	A_13
ATOM	888	N	ASP	99	67.817	45.924	16.394	1.00 21.68	A_13
MOTA	890	ÇA	ASP	99	66.567	46.476	16.886	1.00 10.00	A_13
ATOM	891	CB	ASP	99	66.039	45.604	18.010	1.00 10.00	A_13
MOTA	892	CG	ASP	99	64.648	45.998	18.473	1.00 14.00	A_13
ATOM	893	OD1	ASP	99	64.104	45.272	19.329	1.00 15.19	A_13
ATOM	894	OD2		99	64.089	47.011			
							18.001	1.00 17.01	A_13
ATOM	895	С	ASP	99	66.817	47.871	17.391	1.00 13.06	A_13
ATOM .	896	0	ASP	99	67.528	48.056	18.374	1.00 10.00	A_13
						_			W-13
ATOM	897	N	ASP	100	66.203	48.856	16.746	1.00 15.56	A_13
ATOM	899	CA	ASP	100	66.397	50.232	17.177	1.00 18.23	A_13
MOTA	900	CB	ASP	100	66.121	51.228	16.041	1.00 15.05	
						31.220			A_13
ATOM	901	CG	ASP	100	67.275	52.180	15.838	1.00 11.67	A_13
MOTA	902	נמס	ASP	100	67.602	52.516	14.683	1.00 21.07	A_13
									7-43
ATOM	903		ASP	100	67.879	52.569	16.860	1.00 14.72	A_13
ATOM	904	С	ASP	100	65.610	50.572	18.445	1.00 10.00	A_13
ATOM	905	ō	ASP	100					
					65.767	51.635	19.009	1.00 17.18	A_13
MOTA	906	N	ASP	101	64.755	49.669	18.895	1.00 14.57	A_13
ATOM	908	CA	ASP	101	64.031	49.924			
							20.123	1.00 17.59	A_13
MOTA	909	CB	ASP	101	62,769	49.051	20.236	1.00 12.50	A_13
ATOM	910	CG	ASP	101	61.532	49.721	19.606	1.00 17.12	A_13
MOTA	911		ASP	101	60.599	49.023	19.179	1.00 10.39	A_13
ATOM	912	OD2	ASP	101	61.480	50.962	19.536	1.00 18.09	A_13
MOTA	913	C	ASP	101	64.994	49.766	21.306	1.00 19.33	A_13
MOTA	914	0	ASP	101	64.610	49.972	22.456	1.00 10.00	A_13
ATOM	915		GLU	102					
		N		-	66.213	49.301	21.019	1.00 16.15	A_13
ATOM	917	CA	GLU	102	67.267	49.194	22.044	1.00 13.43	A_13
MOTA	918	CB	GLU	102	68.264	48.085	21.720		X 1 3
								1.00 18.25	A_13
MOTA	919	CG	GLU	102	67.697	46.704	21.636	1.00 10.00	A_13
MOTA	920	CD	GLU	102	66.650	46.467	22.672	1.00 11.18	A_13
MOTA	921				66 070				
			GLU	102	66.872	46.746	23.870	1.00 16.09	A_13
ATOM	922	OE2	GLU	102	65.572	46.033	22.271	1.00 26.76	A_13
ATOM	923	C	GLU	102	68.070	50:495			
							22.007	1.00 11.07	A_13
MOTA	924	0	GLŲ	102	68.103	51.161	20.971	1.00 13.97	A_13
						•	· · · · <del>-</del>		

ATOM	925	N	THR	103	68.774	50.823	23.091	1.00 22.82	A_13
ATOM	927		THR	103	69.606	52.034	23.102	1.00 13.45	A_13
	928		THR	103	69.571				
ATOM						52.793	24.459	1.00 20.78	A_13
ATOM	929		THR	103	68.236	53.228	24.745	1.00 10.69	A_13
ATOM	931		THR	103	70.445	54.046	24.378	1.00 19.45	A_13
MOTA	932	С	THR	103	71.030	51.571	22.822	1.00 12.42	A_13
ATOM	933	0	THR	103	71.639	50.896	23.642	1.00 19.81	A_13
ATOM	934		TRP	104	71.525	51.854	21.626	1.00 10.00	A_13
					72.873				
ATOM	936		TRP	104		51.448	21.248	1.00 13.61	A_13
MOTA	937		TRP	104	72.943	51.221	19.739	1.00 29.21	A_13
ATOM	938	CG	TRP	104	71.970	50.174	19.313	1.00 21.39	A_13
MOTA	939	CD2	TRP	104	72.101	48.760	19.501	1.00 25.13	A_13
ATOM	940		TRP	104	70.937	48.156	18.964	1.00 28.84	A_13
ATOM	941		TRP	104	73.088	47.941			
							20.070	1.00 13.36	A_13
MOTA	942	CD1		104	70.765	50.372	18.694	1.00 21.59	A_13
MOTA	943	NE1	TRP	104	70.139	49.163	18.484	1.00 19.91	A_13
MOTA	945	CZ2	TRP	104	70.738	46.768	18.977	1.00 10.00	A_13
MOTA	946	CZ3	TRP	104	72.888	46.568	20.084	1.00 14.54	A_13
ATOM	947		TRP	104	71.720	45.995	19.539	1.00 11.93	A_13
ATOM	948	c	TRP	104	73.912	52.453	21.725	1.00 16.59	A_13
ATOM	949	0	TRP	104	73.707	53.671	21.642	1.00 12.90	A_13
ATOM	950	N	THR	105	75.013	51.949	22.268	1.00 20.85	A_13
ATOM	952	CA	THR	105	76.040	52.831	22.794	1.00 12.38	A_13
ATOM	953	CB	THR	105	75.974	52.890	24.322	1.00 14.39	A_13
ATOM	954	OG1		105	76.345	51.609	24.849	1.00 16.42	A_13
MOTA	956		THR	105	74.575				A_13
						53.273	24.797	1.00 12.17	A_13
MOTA	957	С	THR	105	77.437	52.378	22.457	1.00 10.00	A_13
MOTA	958	0	THR	105	77.644	51.261	22.012	1.00 18.98	A_13
MOTA	959	N	SER	106	78.385	53.277	22.704	1.00 26.01	A_13
MOTA	961	CA	SER	106	79.809	53.043	22.502	1.00 17.80	A_13
ATOM	962	CB	SER	106	80.466	54.284	21.888	1.00 20.63	A_13
									W_13
MOTA	963	OG	SER	106	79.744	54.756	20.763	1.00 38.89	A_13
MOTA	965	С	SER	106	80.435	52.779	23.880	1.00 34.75	A_13
ATOM	966	0	SER	106	81.652	52.884	24.042	1.00 33.01	A_13
MOTA	967	N	SER	107	79.590	52.494	24.875	1.00 25.87	A_13
ATOM	969	CA	SER	107	80.032	52.221	26.240	1.00 19.68	A_13
ATOM	970	CB	SER	107	80.082	53.510	27.061	1.00 23.47	A_13
							27.001		
MOTA	971	OG	SER	107	78.819	54.158	27.096	1.00 33.70	A_13
MOTA	973	С	SER	107	79.100	51.200	26.892	1.00 13.60	A_13
ATOM	974	0	SER	107	78.460	50.418	26.193	1.00 16.40	A_13
ATOM	975	N	SER	108	79.028	51.205	28.221	1.00 17.31	A_13
ATOM	977	CA	SER	108	78.188	50.259	28.949	1.00 20.12	A_13
ATOM	978	CB	SER	108	78.745	50.009	30.364	1.00 22.63	A_13
MOTA	979								A_13
		OG	SER	108	78.444	51.061	31.271	1.00 27.69	A_13
ATOM	981	С	SER	108	76.702	50.606	29.076	1.00 19.98	A_13
ATOM	982	0	SER	108	75.921	49.785	29.562	1.00 35.96	A_13
ATOM	983	N	LYS	109	76.311	51.820	28.713	1.00 16.24	A_13
MOTA	985	CA	LYS	109	74.907	52.186	28.847	1.00 11.10	A_13
ATOM	986	CB	LYS	109	74.740	53.688	28.690	1.00 12.41	A_13
ATOM	987	CG	LYS	109	73.555				
						54.239	29.462	1.00 32.67	A_13
ATOM	988	CD	LYS	109	73.353	55.732	29.258	1.00 25.94	A_13
ATOM	989	CE	LYS	109	74.535	56.599	29.749	1.00 25.11	A_13
MOTA	990	NZ	LYS	109	74.225	58.070	29.636	1.00 22.70	A_13
MOTA	994	С	LYS	109	74.138	51.424	27.773	1.00 21.67	A_13
MOTA	995	0	LYS	109	74.667	51.210	26.694	1.00 32.76	A_13
ATOM	996	N	GLY	110	72.932	50.955	28.081	1.00 29.60	A_13
ATOM	998	CA	GLY	110	72.156	50.206	27.096		A_13
ATOM	999							1.00 10.31	A_13
		ç	GLY	110	72.965	49.043	26.542	1.00 20.08	A_13
ATOM	1000	0	GLY	110	73.672	48.362	27.285	1.00 11.17	A_13
ATOM	1001	N	TYR	111	72.924	48.859	25.227	1.00 12.05	A_13
ATOM	1003	CA	TYR	111	73.665	47.791	24.583	1.00 13.45	A_13
MOTA	1004	CB	TYR	111	72.713	46.871	23.806	1.00 21.16	A_13
ATOM	1005	ĊĠ	TYR	111	71.776				
ATOM	1005		TYR	111		46.101	24.716	1.00 12.28	A_13
					70.455	46.510	24.906	1.00 14.85	A_13
ATOM	1007	CE1		111	69.618	45.837	25.795	1.00 19.08	A_13
MOTA	1008		TYR	111	72.232	44.995	25.435	1.00 21.86	A_13
MOTA	1009	CE2	TYR	111	71.405	44.314	26.324	1.00 10.00	A_13
MOTA	1010	CZ	TYR	111	70.101	44.740	26.505	1.00 18.51	A_13
ATOM	1011	OH	TYR	111	69.282	44.077	27.398		V-13
MOTA	1013	C	TYR	111				1.00 14.32	.A_13
					74.779	48.335	23.695	1.00 16.73	A_13
ATOM	1014	0	TYR	111	74.540	49.105	22.764	1.00 11.98	A_13
MOTA	1015	N	ASN	112	76.008	47.930	23.999	1.00 11.80	A_13
ATOM	1017	CA	ASN	112	77.184	48.357	23.240	1.00 16.37	A_13
ATOM	1018	CB	ASN	112	78.453	47.867	23.927	1.00 27.52	A_13
MOTA	1019	CG	ASN	112	79.701	48.460	23.324	1.00 20.16	A_13
MOTA	1020		ASN	112	80.327				
MOTA	1021			112		47.861	22.447	1.00 20.99	A_13
*** OI1	1021	NUZ	ASN	114	80.082	49.640	23.801	1.00 15.12	A_13

ATOM	1024	С	ASN	112	77.137	47.809	21.813	1.00 18.08	2 12
ATOM	1025	0	ASN	112	77.288	46.606	21.592	1.00 12.69	A_13
MOTA	1026	N	LEU	113	76.972				A_13
ATOM	1028					48.700	20.844	1.00 11.15	A_13
		CA	LEU	113	76.878	48.296	19.461	1.00 10.00	A_13
ATOM	1029	CB	LEU	113	76.718	49.526	18.568	1.00 10.24	A_13
ATOM	1030	CG	LEU	113	76.325	49.262	17.106	1.00 15.67	A_13
MOTA	1031	CD1	LEU	113	75.155	48.296	17.050	1.00 26.54	A_13
ATOM	1032		LEU	113	75.967	50.555	16.415	1.00 15.60	
MOTA	1033	c	LEU	113	78.037				A_13
						47.403	18.986	1.00 25.17	A_13
ATOM	1034	0	LEU	113	77.799	46.380	18.336	1.00 17.24	A_13
MOTA	1035	N	PHE	114	79.274	47.759	19.327	1.00 28.89	A_13
MOTA	1037	CA	PHE	114	80.442	46.974	18.910	1.00 19.15	A_13
ATOM	1038	CB	PHE	114	81.753	47.579	19.434	1.00 14.60	A_13
ATOM	1039	CG	PHE	114	82.923	46.627	19.374		
ATOM	1040		PHE	114				1.00 18.53	A_13
					83.419	46.175	18.144	1.00 26.13	A_13
ATOM	1041		PHE	114	83.514	46.162	20.547	1.00 17.22	A_13
ATOM	1042		PHE	114	84.475	45.271	18.086	1.00 10.43	A_13
MOTA	1043	CE2	PHE	114	84.571	45.259	20.502	1.00 16.51	A_13
MOTA	1044	CZ	PHE	114	85.052	44.815	19.260	1.00 15.54	A_13
ATOM	1045	C	PHE	114	80.359	45.508	19.306	1.00 10.00	
MOTA	1046	ŏ	PHE	114					A_13
ATOM					80.437	44.625	18.445	1.00 33.07	A_13
	1047	N	LEU	115	80.206	45.249	20.600	1.00 12.18	A_13
ATOM	1049	CA	LEU	115	80.113	43.877	21.103	1.00 10.59	A_13
MOTA	1050	CB	LEU	115	79.874	43.895	22.616	1.00 14.14	Á_13
MOTA	1051	CG	LEU	115	81.082	43.937	23.578	1.00 34.39	A_13
ATOM	1052	CD1	LEU	115	82.337	44.354	22.863	1.00 14.93	
ATOM	1053		LEU	115					A_13
					80.815	44.836	24.793	1.00 13.42	A_13
ATOM	1054	C	LEU	115	79.019	43.080	20.379	1.00 12.06	A_13
MOTA	1055	0	LEU	115	79.298	42.109	19.675	1.00 13.35	A_13
MOTA	1056	N	VAL	116	77.786	43.558	20.459	1.00 13.11	A_13
MOTA	1058	CA	VAL	116	76.678	42.875	19.814	1.00 12.97	
ATOM	1059	СВ	VAL	116	75.343	43.569			A_13
ATOM	1060	_	VAL				20.129	1.00 28.07	A_13
				116	74.200	42.926	19.340	1.00 17.32	A_13
MOTA	1061		VAL	116	75.074	43.491	21.617	1.00 22.14	A_13
MOTA	1062	С	VAL	116	76.862	42.724	18.313	1.00 10.00	A_13
MOTA	1063	0	VAL	116	76.473	41.716	17.755	1.00 14.68	A_13
ATOM	1064	N	ALA	117	77.481	43.706	17.667	1.00 10.80	A_13
ATOM	1066	CA	ALA	117	77.726	43.662			
ATOM	1067	CB	ALA				16.224	1.00 18.28	A_13
				117	78.223	45.014	15.727	1.00 14.94	A_13
MOTA	1068	C	ALA	117	78.735	42.579	15.863	1.00 25.24	A_13
MOTA	1069	0	ALA	117	78.562	41.872	14.861	1.00 18.50	A_13
ATOM	1070	N	ALA	118	79.795	42.458	16.665	1.00 24.40	A_13
ATOM	1072	CA	ALA	118	80.829	41.451	16.422	1.00 11.80	A_13
ATOM	1073	CB	ALA	118	81.945	41.590		1.00 11.00	W_13
ATOM	1074		ALA				17.447	1.00 19.28	A_13
		C		118	80.178	40.056	16.496	1.00 10.00	A_13
MOTA	1075	0	ALA	118	80.426	39.183	15.660	1.00 10.00	A_13
ATOM	1076	N	HIS	119	79.309	39.875	17.487	1.00 19.01	A_13
MOTA	1078	CA	HIS	119	78.587	38.624	17.674	1.00 14.36	A_13
ATOM	1079	CB	HIS	119	77.725	38.751	18.924	1.00 10.00	A_13
ATOM	1080	CG	HIS	119	. 76.796	37.602	19.166	1.00 10.00	
ATOM	1081		HIS	119	75.691				A_13
ATOM						37.187	18.498	1.00 14.94	A_13
	1082		HIS	119	76.905	36.783	20.263	1.00 20.37	A_13
ATOM	1084	CEI	HIS	119	75.917	35.909	20.270	1.00 17.53	A_13
MOTA	1085	NE2	HIS	119	75.161	36.134	19.208	1.00 17.55	A_13
MOTA	1086	С	HIS	119	77.741	38.339	16.419	1.00 10.00	A_13
ATOM	1087	0	HIS	119	77.779	37.245	15.856	1.00 10.64	A_13
ATOM	1088	N	GLU	120	77.004	39.343	15.968		
ATOM	1090	CA	GĽÜ	120	76.174			1.00 22.95	A_13
ATOM	1091					39.224	14.775	1.00 23.96	A_13
		CB	GLU	120	75.429	40.545	14.502	1.00 17.19	A_13
MOTA	1092	CG	GLU	120	74.373	40.889	15.555	1.00 16.14	A_13
MOTA	1093	CD	GLU	120	73.492	39.691	15.929	1.00 10.00	A_13
ATOM	1094	OE1	GLU	120	73.478	39.354	17.122	1.00 17.94	A_13
MOTA	1095		GLU	120	72.844	39.078	15.047		
ATOM	1096	c	GLU	120	76.992		13 540	1.00 17.03	A_13
ATOM	1097					38.832	13.549	1.00 11.45	A_13
		0	GLU	120	76.594	37.946	12.772	1.00 13.34	A_13
ATOM	1098	N	PHE	121	78.127	39.498	13.353	1.00 10.00	A_13
MOTA	1100	CA	PHE	121	78.959	39.187	12.216	1.00 14.70	A_13
MOTA	1101	CB	PHE	121	80.040	40.245	12.039	1.00 10.00	A_13
ATOM	1102	CG	PHE	121	79.481	41.623	11.792	1.00 21.57	A_13
ATOM	1103		PHE	121	80.235	42.764			
MOTA	1104		PHE	121	78.164		12.069	1.00 16.73	A_13
ATOM	1105					41.788	11.331	1.00 13.91	A_13
			PHE	121	79.682	44.054	11.891	1.00 11.69	A_13
ATOM	1106	CE2		121	77.615	43.066	11.152	1.00 18.93	A_13
MOTA	1107	CZ	PHE	121	78.373	44.192	11.436	1.00 10.00	A_13
MOTA	1108	C	PHE	121	79.505	37.756	12.283	1.00 17.14	A_13
ATOM	1109	0	PHE	121	79.642	37.104	11.256	1.00 13.04	A_13
MOTA	1110	N	GLY	122	79.738	37.245	13.490		
					13.135	37.243	13.450	1.00 16.60	A_13

					•				
ATOM	1112	CA	GLY	122	00 202	25 072	13.627	1 00 10 45	
					80.202	35.872		1.00 19.45	A_13
ATOM	1113		GLY	122	79.162	34.982	12.966	1.00 18.55	A_13
MOTA	1114	0	GLY	122	79.500	33.988	12.306	1.00 10.03	A_13
ATOM	1115	N	HIS	123	77.892	35.361	13.140	1.00 18.22	A_13
ATOM	1117		HIS	123	76.753	34.665	12.525	1.00 16.31	
ATOM	1118								A_13
			HIS	123	75.424	35.224	13.031	1.00 11.35	A_13
MOTA	1119	CG	HIS	123	75.049	34.768	14.403	1.00 10.33	A_13
ATOM	1120	CD2	HIS	123	74.552	35.454	15.457	1.00 16.64	A_13
MOTA	1121	ND1	HTC	123	75.097	33.450	14.782	1.00 18.04	
									A_13
MOTA	1123	CE1		123	74.638	33.332	16.017	1.00 16.66	A_13
MOTA	1124	NE2	HIS	123	74.301	34.533	16.450	1.00 25.32	A_13
MOTA	1125	С	HIS	123	76.771	34.853	10.997	1.00 13.66	A_13
MOTA	1126		HIS	123	76.565	33.901	10.246	1.00 10.82	
									A_13
MOTA	1127	N	SER	124	77.006	36.082	10.539	1.00 13.57	A_13
MOTA	1129	CA	SER	124	77.030	36.368	9.099	1.00 12.03	A_13
MOTA	1130	CB	SER	124	77.311	37.863	8.832	1.00 10.35	A_13
MOTA	1131	OG	SER	124	76.399	38.706	9.510	1.00 14.26	A_13
ATOM	1133	C	SER	124	78.117	35.548			V-13
							8.422	1.00 21.45	A_13
MOTA	1134	0	SER	124	78.079	35.333	7.210	1.00 10.00	A_13
MOTA	1135	N	LEU	125	79.091	35.108	9.216	1.00 10.00	A_13
ATOM	1137	CA	LEU	125	80.222	34.340	8.707	1.00 19.28	A_13
ATOM	1138	CB	LEU	125	81.521	34.754	9.422	1.00 22.39	A_13
ATOM	1139	CG	LEU						A_13
	-			125	81.849	36.258	9.340	1.00 10.00	A_13
MOTA	1140	CD1		125	83.063	36.622	10.190	1.00 10.00	A_13
MOTA	1141	CD2	LEU	125	82.029	36.651	7.873	1.00 10.00	A_13
ATOM	1142	С	LEU	125	79.986	32.851	8.843	1.00 10.00	A_13
MOTA	1143	ō	LEU	125	80.759	32.056	8.329	1.00 23.27	2-73
									A_13
MOTA	1144	N	GLY	126	78.932	32.477	9.563	1.00 22.87	A_13
MOTA	1146	CA	GLY	126	78.604	31.070	9.720	1.00 17.27	A_13
MOTA	1147	C	GLY	126	78.781	30.464	11.094	1.00 11.71	A_13
MOTA	1148	Ó	GLY	126	78.784	29.244	11.236	1.00 24.16	
MOTA									A_13
	1149	N	LEU	127	78.972	31.297	12.105	1.00 18.95	A_13
MOTA	1151	CA	LEU	127	79.152	30.790	13.457	1.00 22.84	A_13
MOTA	1152	CB	LEU	127	80.113	31.693	14.252	1.00 11.92	A_13
MOTA	1153	CG	LEU	127	81.244	30.969	14.983	1.00 18.83	A_13
MOTA	1154		LEU	127					
					82.096	30.197	13.979	1.00 16.63	A_13
MOTA	1155		LEU	127	82.104	31.970	15.760	1.00 22.15	A_13
MOTA	1156	С	LEU	127	77.802	30.699	14.163	1.00 21.02	A_13
ATOM	1157	0	LEU	127	76.996	31.629	14.098	1.00 14.68	A_13
ATOM	1158	N	ASP	128	77.563	29.572			
							14.828	1.00 18.87	A_13
MOTA	1160	CA	ASP	128	76.336	29.345	15.571	1.00 16.46	A_13
MOTA	1161	CB	ASP	128	75.996	27.855	15.540	1.00 17.60	A_13
ATOM	1162	CG	ASP	128	74.577	27.552	15.996	1.00 23.55	A_13
MOTA	1163	OD1	ASP	128	73.796	28.488	16.258	1.00 10.00	A_13
ATOM	1164		ASP	128	74.236	26.355			
							16.087	1.00 32.36	A_13
ATOM	1165	C	ASP	128	76.634	29.803	16.995	1.00 10.00	A_13
ATOM	1166	0	asp	128	77.650	30.420	17.244	1.00 29.54	A_13
ATOM	1167	N	HIS	129	75.714	29.565	17.912	1.00 10.00	A_13
MOTA	1169	CA	HIS	129	75.910	29.955	19.289	1.00 10.00	A 13
ATOM	1170	CB	HIS	129	74.582	30.033	20.029	1.00 21.30	A_13
ATOM	1171	CG	HIS	129					W_13
					73.798	31.282	19.761	1.00 24.16	A_13
MOTA	1172		HIS	129	74.180	32.585	19.725	1.00 10.00	A_13
MOTA	1173	ND1	HIS	129	72.460	31.263	19.476	1.00 21.70	A_13
ATOM	1175	CE1	HIS	129	72.031	32.501	19.271	1.00 10.27	A_13
ATOM	1176	NE2	HIS	129	73.057	33.319	19.407	1.00 14.37	A_13
MOTA	1177	C	HIS	129	76.780				
ATOM	1178	ŏ	HIS	129		28.947	19.992	1.00 30.04	A_13
					76.624	27.730	19.822	1.00 22.13	A_13
ATOM	1179	N	SER	130	77.628	29.468	20.860	1.00 18.60	A_13
ATOM	1181	CA	SER	130	78.534	28.662	21.636	1.00 10.79	. A_13
MOTA	1182	CB	SER	130	79.849	29.435	21.816	1.00 21.31	A_13
MOTA	1183	ŌĞ	SER	130	80.782				
						28.731	22.616	1.00 16.34	A_13
ATOM	1185	C	SER	130	77.898	28.368	22.987	1.00 31.13	A_13
ATOM	1186	0	SER	130	76.962	29.060	23.440	1.00 15.87	A_13
MOTA	1187	N	LYS	131	78.402	27.319	23.619	1.00 13.13	A_13
ATOM	1189	CA	LYS	131	77.924	26.925	24.928	1.00 13.21	A_13
ATOM	1190								A_13
		CB	LYS	131	77.656	25.414	24.990	1.00 18.85	A_13
MOTA	1191	CG	LYS	131	78.689	24.541	24.303	1.00 32.55	A_13
ATOM	1192	CD	LYS	131	78.547	24.601	22.790	1.00 41.54	A_13
MOTA	1193	CE	LYS	131	79.909	24.672	22.117	1.00 19.64	A_13
ATOM	1194	NZ	LYS	131					
					80.747	25.799	22.617	1.00 13.47	A_13
ATOM	1198	C	LYS	131	78.922	27.379	25.982	1.00 10.00	A_13
MOTA	1199	0	LYS	131	78.666	27.260	27.185	1.00 13.35	A_13
ATOM	1200	N	ASP	132	80.025	27.968	25.519	1.00 13.47	A_13
ATOM	1202	CA	ASP	132	81.097	28.487	26.375		A_13
ATOM	1203	CB	ASP	132				1.00 10.04	ú-13
MOTA					82.376	28.617	25.522	1.00 18.14	A_13
	1204	CG	ASP	132	83.649	28.821	26.345	1.00 16.54	A_13
MOTA	1205	OD1	ASP	132	84.645	28.132	26.028	1.00 36.08	A_13

MOTA	1206	OD2	ASP	132	83.685	29.660	27.276	1.00 15.60	1 12
ATOM	1207	C	ASP	132					A_13
					80.603	29.875	26.836	1.00 18.74	A_13
ATOM	1208	0	ASP	132	80.559	30.816	26.038	1.00 14.61	A_13
ATOM	1209	N	PRO	133	80.305	30.039	28.142	1.00 15.61	A_13
ATOM	1210	CD	PRO	133	80.617	29.127	29.251	1.00 21.19	
MOTA	1211	CA	PRO	133	79.818				A_13
						31.320	28.662	1.00 10.00	A_13
MOTA	1212	CB	PRO	133	79.542	31.007	30.135	1.00 10.00	A_13
ATOM	1213	CG	PRO	133	80.633	30.063	30.450	1.00 30.94	A_13
MOTA	1214	С	PRO	133	80.834	32.444			
							28.511	1.00 22.87	A_13
MOTA	1215	0	PRO	133	80.526	33.574	28.742	1.00 21.65	A_13
MOTA	1216	N	GLY	134	82.070	32.115	28.174	1.00 20.95	A_13
ATOM	1218	CA	GLY	134	83.055	33.167	28.028	1.00 15.22	
ATOM	1219	C	GLY	134					A_13
					83.182	33.578	26.581	1.00 34.54	A_13
ATOM	1220	0	GLY	134	83.962	34.488	26.252	1.00 18.06	A_13
ATOM	1221	N	ALA	135	82.490	32.846	25.706	1.00 21.09	A_13
ATOM	1223	CA	ALA	135		22.030			W_13
					82.547	33.110	24.263	1.00 27.50	A_13
ATOM	1224	CB	ALA	135	82.131	31.858	23.453	1.00 10.00	A_13
ATOM	1225	С	ALA	135	81.722	34.308	23.814	1.00 21.74	A_13
ATOM	1226	0	ALA	135 .	80.641	34.556	24.328		
ATOM	1227	N						1.00 13.84	A_13
			LEU	136	82.220	34.990	22.787	1.00 19.10	A_13
MOTA	1229	CA	LEU	136	81.540	36.140	22.203	1.00 21.65	A_13
ATOM	1230	CB	LEU	136 .	82.448	36.803	21.161	1.00 10.00	A_13
ATOM	1231	CG	LEU	136	81.964	37.898	20.201	1.00 17.22	7_13
ATOM	1232		LEU	136					A_13
					81.250	37.296	19.024	1.00 24.18	A_13
ATOM	1233		LEU	136	81.113	38.896	20.905	1.00 10.00	A_13
ATOM	1234	C	LEU	136	80.250	35.632	21.558	1.00 19.32	A_13
MOTA	1235	0	LEU	136	79.266	36.359			7_13
MOTA							21.458	1.00 26.20	A_13
	1236	N	MET	137	80.297	34.409	21.029	1.00 10.00	A_13
MOTA	1238	CA	MET	137	79.123	33.791	20.423	1.00 10.02	A_13
MOTA	1239	CB	MET	137	79.507	32.691	19.428	1.00 15.14	
ATOM	1240	CG	MET	137					A_13
					80.181	33.223	18.169	1.00 16.42	A_13
ATOM	1241	SD	MET	137	79.366	34.665	17.397	1.00 10.65	A_13
ATOM	1242	CE	MET	137	77.848	34.005	16.975	1.00 10.87	A_13
ATOM	1243	С	MET	137	78.122	33.256	21.447	1.00 12.70	7-13
MOTA	1244	ō	MET						A_13
				137	77.187	32.539	21.087	1.00 10.00	A_13
MOTA	1245	N	PHE	138	78.295	33.627	22.713	1.00 18.70	A_13
MOTA	1247	CA	PHE	138	77.370	33.196	23.759	1.00 24.08	A_13
MOTA	1248	CB	PHE	138	77.954				
ATOM						33.448	25.159	1.00 24.15	A_13
	1249	CG	PHE	138	77.306	32.617	26.240	1.00 29.38	A_13
MOTA	1250	CD1	PHE	138	76.694	33.222	27.336	1.00 27.07	A_13
ATOM	1251	CD2	PHE	138	77.253	31.226	26.123	1.00 21.37	
MOTA	1252		PHE	138					A_13
					76.033	32.455	28.289	1.00 30.35	A_13
MOTA	1253		PHE	138	76.599	30.458	27.065	1.00 19.58	A_13
ATOM	1254	CZ	PHE	138	75.986	31.070	28.154	1.00 17.69	A_13
MOTA	1255	C	PHE	138	76.074	33.992			
ATOM	1256						23.513	1.00 14.20	A_13
		0	PHE	138	76.115	35.105	23.014	1.00 10.27	A_13
ATOM	1257	N	PRO	139	74.899	33.366	23.730	1.00 13.04	A_13
ATOM	1258	CD	PRO	139	74.664	31.975	24.131	1.00 11.17	A_13
ATOM	1259	CA	PRO	139	73.619	34.043	23.504		2-43
MOTA	1260	CB	PRO	139				1.00 18.27	A_13
					72.625	32.875	23.384	1.00 14.33	` A_13
ATOM	1261	CG	PRO	139	73.474	31.634	23.305	1.00 24.22	A_13
MOTA	1262	Ç	PRO	139	73.162	35.018	24.584	1.00 16.51	A_13
ATOM	1263	0	PRO	139	72.023	35.467			
MOTA	1264	N	ILE	140	74.034		24.535	1.00 24.45	A_13
						35.375	25.524	1.00 23.16	A_13
MOTA	1266	CA	ILE	140	73.652	36.290	26.604	1.00 25.00	A_13
MOTA	1267	CB	ILE	140	73.688	35.559	27.966	1.00 12.10	A_13
ATOM	1268	CG2	ILE	140	73.336	36.519	29.085	1.00 12.62	A_13
MOTA	1269		ILE	140			27.003	1.00 12.02	W_13
					72.738	34.341	27.904	1.00 22.67	A_13
MOTA	1270		ILE	140	72.827	33.353	29.073	1.00 27.73	A_13
MOTA	1271	С	ILE	140	74.584	37.489	26.621	1.00 30.64	A_13
ATOM	1272	0	ILE	140	75.778	37.317	26.682		7_13
MOTA	1273	N	TYR	141				1.00 23.16	A_13
					74.033	38.694	26.532	1.00 21.05	A_13
MOTA	1275	CA	TYR	141	74.851	39.901	26.528	1.00 20.10	A_13
MOTA	1276	CB	TYR	141	74.017	41.122	26.129	1.00 17.66	A_13
ATOM	1277	CG	TYR	141	74.784	42.433			W_T3
ATOM	1278						26.103	1.00 22.24	A_13
			TYR	141	74.711	43.318	27.171	1.00 18.07	A_13
MOTA	1279		TYR	141	75.386	44.527	27.144	1.00 19.84	A_13
ATOM	1280	CD2	TYR	141	75.563	42.798	24.999	1.00 18.08	A_13
ATOM	1281		TYR	141	76.244				W_13
						44.008	24.961	1.00 10.00	A_13
ATOM	1282	CZ	TYR	141	76.149	44.867	26.038	1.00 25.17	A_13
MOTA	1283	OH	TYR	141	76.814	46.070	26.043	1.00 30.78	A_13
MOTA	1285	С	TYR	141	75.533	40.169	27.852		7 12
MOTA	1286	ŏ	TYR	141				1.00 19.61	A_13
					74.910	40.146	28.913	1.00 16.08	A_13
ATOM	1287	N	THR	142	76.817	40.476	27.772	1.00 26.26	A_13
MOTA	1289	CA	THR	142	77.612	40.788	28.944	1.00 24.52	A_13
ATOM	1290	CB	THR	142	78.498				
ATOM						39.568	29.362	1.00 10.00	A_13
2104	1291	T	THR	142	77.664	38.587	29.981	1.00 37.30	A_13

MOTA	1293	CG2	THR	142	79.543	39.961	30.390	1.00 14.88	A_13
MOTA	1294	C	THR	142	78.467	41.976	28.580	1.00 25.46	A_13
MOTA MOTA	1295 1296	О И	THR TYR	142 143	78.980 78.575	42.058 42.947	27.464 29.476	1.00 10.00 1.00 20.23	A_13
ATOM	1298	CA	TYR	143	79.412	44.079	29.133	1.00 20.23	A_13 A_13
MOTA	1299	СВ	TYR	143	79.024	45.363	29.854	1.00 35.01	A_13
MOTA	1300	CG	TYR	143	79.834	46.531	29.347	1.00 16.01	A_13
MOTA	1301	CD1		143	79.776	46.910	27.998	1.00 12.56	A_13
ATOM	1302	CE1		143	80.554	47.961	27.510	1.00 19.23	A_13
MOTA MOTA	1303 1304	CD2 CE2	TYR TYR	143 143	80.690 81.478	47.230	30.196	1.00 19.43	A_13
ATOM	1304	CZ	TYR	143	81.403	48.287 48.643	29.719 28.376	1.00 15.52 1.00 12.56	A_13 A_13
ATOM	1306	ОН	TYR	143	82.193	49.654	27.892	1.00 18.85	A_13
MOTA	1308	C	TYR	143	80.871	43.754	29.382	1.00 25.10	A_13
MOTA	1309	0	TYR	143	81.373	43.846	30.503	1.00 28.90	A_13
MOTA	1310	N	THR	144	81.539	43.375	28.303	1.00 35.25	A_13
MOTA	1312	CA	THR	144	82.946	43.029	28.336	1.00 38.86	A_13
ATOM ATOM	1313 1314	CB	THR THR	144 144	83.158 82.129	41.568 41.219	27.873 26.934	1.00 23.22 1.00 35.22	A_13 A_13
MOTA	1316		THR	144	83.105	40.616	29.082	1.00 33.22	A_13
ATOM	1317	C	THR	144	83.720	44.017	27.488	1.00 21.63	A_13
MOTA	1318	0	THR	144	84.434	43.651	26.556	1.00 37.44	A_13
MOTA	1319	N	GLY	145	83.504	45.288	27.798	1.00 14.47	A_13
MOTA	1321	CA	GLY	145	84.200	46.375	27.131	1.00 24.39	A_13
MOTA MOTA	1322 1323	0	GLY GLY	145 145	84.119 84.053	46.536 45.565	25.628 24.877	1.00 41.65 1.00 42.39	A_13 A_13
MOTA	1324	N	LYS	146	84.122	47.792	25.195	1.00 42.39	A_13
ATOM	1326	CA	LYS	146	84.059	48.103	23.778	1.00 29.29	A_13
ATOM	1327	CB	LYS	146	83.260	49.392	23.539	1.00 26.47	A_13
ATOM	1328	CG	LYS	146	83.087	49.721	22.059	1.00 33.24	A_13
MOTA	1329	CD	LYS	146	82.812	51.194	21.833	1.00 13.70	A_13
ATOM ATOM	1330 1331	CE NZ	LYS LYS	146 146	82.620 83.766	51.497 51.122	20.343 19.477	1.00 18.35 1.00 30.66	A_13
MOTA	1335	C	LYS	146	85.491	48.297	23.308	1.00 30.66	A_13 A_13
ATOM	1336	ō	LYS	146	86.028	49.412	23.382	1.00 46.44	A_13
ATOM	1337	N	SER	147	86.130	47.206	22.898	1.00 34.67	A_13
MOTA	1339	CA	SER	147	87.509	47.258	22.416	1.00 30.76	A_13
MOTA	1340	CB	SER	147	87.624	48.258	21.249	1.00 24.56	A_13
MOTA MOTA	1341 1343	OG C	SER SER	147 147	86.638	48.002	20.257	1.00 31.81	A_13
ATOM	1344	ō	SER	147	88.464 88.789	47.626 48.806	23.567 23.789	1.00 33.60 1.00 39.96	A_13 A_13
ATOM	1345	N	HIS	148	88.862	46.611	24.331	1.00 36.71	A_13
ATOM	1347	CA	HIS	148	89.778	46.769	25.467	1.00 34.40	A_13
ATOM	1348	CB	HIS	148	89.307	47.862	26.438	1.00 26.40	A_13
MOTA	1349	CG	HIS	148	90.251	49.022	26.537	1.00 39.11	A_13
MOTA MOTA	1350 1351		HIS HIS	148 148	90.929	49.542	27.588	1.00 30.52	A_13
ATOM	1353		HIS	148	90.635 91.511	49.767 50.681	25.437 25.807	1.00 37.71 1.00 29.04	A_13 A_13
ATOM	1354		HIS	148	91.707	50.567	27.110	1.00 29.03	A_13
ATOM	1356	С	HIS	148	89.949	45.436	26.190	1.00 39.41	A_13
MOTA	1357	0	HIS	148	90.134	45.373	27.411	1.00 35.01	A_13
MOTA MOTA	1358	N	PHE	149	89.840	44.386	25.383	1.00 25.35	A_13
ATOM	1360 1361	CA CB	PHE PHE	149 149	89.996 88.788	42.966 42.423	25.721 26.495	1.00 30.54 1.00 33.34	A_13
ATOM	1362	CG		149	88.951	42.440	27.996	1.00 33.34	A_13 A_13
MOTA	1363		PHE	149	89.387	41.302	28.673	1.00 30.46	A_13
MOTA	1364		PHE	149	88.624	43.575	28.740	1.00 40.67	A_13
ATOM	1365		PHE	149	89.492	41.293	30.075	1.00 18.92	A_13
MOTA MOTA	1366 1367	CE2	PHE	149 149	88.728 89.161	43.574	30.136	1.00 23.23	A_13
ATOM	1368	C	PHE	149	90.026	42.430 42.366	30.803 24.295	1.00 17.03 1.00 41.76	A_13
ATOM	1369	ŏ	PHE	149	89.967	43.119	23.307	1.00 40.43	A_13 A_13
'ATOM	1370	N	MET	150	90.132	41.050	24.142	1.00 31.30	A_13
ATOM	1372	CA	MET	150	90.152	40.531	22.779	1.00 20.65	A_13
MOTA	1373	CB	MET	150	91.588	40.195	22.352	1.00 28.29	A_13
ATOM	1374	CG	MET	150	92.494	41.436	22.188	1.00 34.71	A_13
ATOM ATOM	1375 1376	SD CE	MET MET	150 150	91.750 92.512	42.780	21.185 19.518	1.00 67.91	A_13
ATOM	1377	C	MET	150	89.201	42.498 39.370	22.497	1.00 22.43 1.00 21.51	A_13 A_13
MOTA	1378	õ	MET	150	88.498	38.901	23.391	1.00 25.37	A_13 A_13
MOTA	1379	N	LEU	151	89.159	38.938	21.240	1.00 13.78	A_13
ATOM	1381	CA	LEU	151	88.313	37.825	20.834	1.00 14.73	A_13
MOTA	1382	CB	LEU	151	88.435	37.589	19.321	1.00 15.49	A_13
ATOM ATOM	1383	CG	LEU	151	87.535	36.511	18.691	1.00 27.05	A_13
MOTA	1384 1385		LEU LEU	151 151	86.070 87.879	36.915 36.310	18.847 17.208	1.00 10.98 1.00 15.73	A_13 A 13
ATOM	1386	C	LEU	151	88.732	36.563	21.600	1.00 25.01	A_13 A_13
		-			-3.732		22.000	2 20.01	

ATOM	1387	0	LEU	151	89.912	36.178	21.589	1.00 17.37	h 12
									A_13
ATOM	1388	N	PRO	152	87.777	35.927	22.306	1.00 10.37	A_13
ATOM	1389	CD	PRO	152	86.425	36.450	22.575	1.00 15.35	A_13
ATOM	1390	CA	PRO	152	88.030	34.712	23.087	1.00 11.49	A_13
ATOM	1391	CB	PRO	152	86.658	34.412	23.702	1.00 15.98	A_13
ATOM	1392	CG	PRO	152	86.083	35.789	23.898	1.00 27.60	A_13
MOTA	1393	С	PRO	152	88.533	33.553	22.230	1.00 18.06	A_13
ATOM	1394	0	PRO	152	88.160	33.430	21.063	1.00 16.21	A_13
MOTA	1395	N	ASP	153	89.350	32.696	22.836	1.00 15.86	A_13
	1397								
MOTA		CA	ASP	153	89.933	31.526	22.185	1.00 20.25	A_13
ATOM	1398	CB	ASP	153	90.632	30.630	23.227	1.00 18.17	A_13
ATOM	1399	CG	ASP	153	91.843	31.301	23.908	1.00 24.01	A_13
MOTA	1400	ODI	ASP	153	92.517	32.159	23.284	1.00 14.96	A_13
ATOM									A_13 .
	1401		ASP	153	92.131	30.937	25.077	1.00 20.20	A_13 '
MOTA	1402	С	ASP	153	88.887	30.678	21.452	1.00 24.64	A_13
MOTA	1403	0	ASP	153	89.113	30.221	20.330	1.00 13.51	A_13
MOTA	1404	N	ASP	154	87.757	30.453	22.114	1.00 24.11	A_13
MOTA	1406	CA	ASP	154	86.664	29.657	21.577	1.00 19.19	A_13
ATOM	1407	CB	ASP	154	85.527	29.632	22.587	1.00 18.27	A_13
ATOM	1408	CG	ASP	154	84.406	28.751	22.161	1.00 24.26	A_13
ATOM	1409	001	ASP	154	83.314	29.291	21.950	1.00 20.97	
									A_13
ATOM	1410	OD2	ASP	154	84.609	27.530	22.031	1.00 20.32	A_13
MOTA	1411	С	ASP	154	86.162	30.170	20.229	1.00 18.99	A_13
MOTA	1412	0	ASP	154	86.043	29.408	19.277	1.00 22.56	A_13
MOTA	1413	N	ASP	155	85.873	31.465	20.158	1.00 16.11	A_13
ATOM	1415	CA	ASP	155					
					85.407	32.078	18.917	1.00 25.30	A_13
MOTA	1416	CB	ASP	155	85.011	33.527	19.158	1.00 13.32	A_13
ATOM	1417	CG	ASP	155	83.975	33.655	20.249	1.00 11.19	A_13
ATOM	1418	OD1	ASP	155	84.347	34.136	21.332	1.00 12.26	A_13
ATOM	1419		ASP	155					
					82.810	33.255	20.029	1.00 10.00	A_13
MOTA	1420	С	ASP	155	86.461	31.992	17.828	1.00 13.98	A_13
ATOM	1421	0	ASP	155	86.141	31.656	16.687	1.00 14.08	A_13
ATOM	1422	N	VAL	156	87.713	32.310	18.160	1.00 16.49	A_13
ATOM	1424	CA	VAL	156					
					88.771	32.201	17.159	1.00 27.34	A_13
MOTA	1425	CB	VAL	156	90.145	32.826	17.625	1.00 23.59	A_13
ATOM	1426	CG1	VAL	156	90.327	32.750	19.119	1.00 13.94	A_13
MOTA	1427	CG2	VAL	156	91.312	32.153	16.919	1.00 21.70	A_13
ATOM	1428								
		C	VAL	156	88.874	30.738	16.657	1.00 16.95	A_13
MOTA	1429	0	VAL	156	88.946	30.506	15.448	1.00 13.79	A_13
MOTA	1430	N	GLN	157	88.762	29.763	17.561	1.00 19.45	A_13
MOTA	1432	CA	GLN	157	88.796	28.352	17.154	1.00 30.53	A_13
ATOM	1433	CB	GLN	157					7-13
					88.579	27.422	18.353	1.00 23.08	A_13
MOTA	1434	CG	GLN	157	89.633	27.521	19.452	1.00 24.83	A_13
MOTA	1435	CD	GLN	157	90.950	26.872	19.089	1.00 20.26	A_13
ATOM	1436	OE1	GLN	157	91.743	27.422	18.316	1.00 25.80	A_13
MOTA	1437		GLN	157	91.204	25.702	19.673	1.00 38.67	A_13
ATOM									
	1440	C	GLN	157	87.667	28.136	16.148	1.00 14.16	A_13
ATOM	1441	0	GLN	157	87.869	27.541	15.096	1.00 14.11	A_13
MOTA	1442	N	GLY	158∙	86.505	28.709	16.437	1.00 19.16	A_13
ATOM	1444	CA	GLY	158	85.361	28.584	15.551	1.00 12.79	A_13
ATOM	1445	c c	GLY	158					
					85.510	29.144	14.143	1.00 24.46	A_13
ATOM	1446	0	GLY	158	85.181	28.449	13.177	1.00 18.77	A_13
MOTA	1447	N	ILE	159	85.936	30.403	13.989	1.00 22.41	A_13
MOTA	1449	CA	ILE	159	86.091	30.946	12.628	1.00 31.18	A_13
ATOM	1450	CB	ILE	159	86.300	32.508	12.532	1.00 23.53	A_13
ATOM	1451		ILE	159					
					84.991	33.203	12.177	1.00 17.28	A_13
MOTA	1452		ILE		87.022	33.063	13.758	1.00 15.28	A_13
MOTA	1453	CD1	ILE	159	88.507	32.949	13.707	1.00 14.71	A_13
MOTA	1454	С	ILE		87.226	30.280	11.875	1.00 10.56	A_13
ATOM	1455	ō	ILE		87.167	30.139	10.653	1.00 18.79	2-13
ATOM	1456								A_13
		N	GLN		88.287	29.927	12.590	1.00 20.71	A_13
MOTA	1458	CA	GLN		89.411	29.294	11.943	1.00 10.00	A_13
MOTA	1459	CB	GLN	160	90.640	29.274	12.855	1.00 10.00	A_13
MOTA	1460	CG	GLN		91.114	30.690	13.182	1.00 13.93	
ATOM	1461					20.030			A_13
		CD	GLN		92.402	30.754	13.981	1.00 25.61	A_13
MOTA	1462		GLN		92.814	29.786	14.629	1.00 19.40	A_13
MOTA	1463	NE2	GLN	160	93.042	31.915	13.950	1.00 24.78	A_13
MOTA	1466	С	GLN		89.000	27.917	11.477	1.00 10.00	7-13
ATOM	1467								A_13
		0	GLN		89.458	27.481	10.432	1.00 21.73	A_13
MOTA	1468	N	SER	161	88.068	27.268	12.186	1.00 10.00	A_13
ATOM	1470	CA	SER	161	87.610	25.946	11.760	1.00 11.63	A_13
ATOM	1471	CB	SER		86.688	25.292	12.800	1.00 18.40	
ATOM	1472	OG							A_13
			SER		85.365	25.795	12.759	1.00 15.44	A_13
ATOM	1474	C	SER		86.913	26.048	10.396	1.00 26.18	A_13
MOTA	1475	0	SER	161	86.839	25.065	9.654	1.00 13.96	A_13
MOTA	1476	N	LEU	162	86.428	27.247	10.070	1.00 19.36	A_13
ATOM	1478	CA	LEU		85.749	27.493			
	2-2.0	VA.	0 ندس		03.749	41.473	8.808	1.00 17.21	A_13

MOTA		CB LEU	162	84.584	28.477	9.007	1.00 14.37	A_13
MOTA		CG LEU	162	83.489	28.144	10.021	1.00 31.09	A_13
MOTA		CD1 LEU	162	82.596	29.351	10.217	1.00 14.96	A_13
ATOM		CD2 LEU	162	82.672	26.949	9.548	1.00 23.87	A_13
ATOM ATOM		C LEU O LEU	162 162	86.654 86.596	28.080 27.680	7.744 6.584	1.00 11.98 1.00 15.25	A_13
ATOM			163	87.459	29.063	8.135		A_13
ATOM		N TYR CA TYR	163	88.320	29.796	7.204	1.00 26.64	A_13
MOTA			163	87.977	31.289	7.277	1.00 18.28 1.00 26.89	A_13
MOTA		CB TYR	163	86.519	31.600	7.039		A_13
MOTA		CD1 TYR	163	86.027	31.744	5.749	1.00 18.80 1.00 10.00	A_13
MOTA		CE1 TYR	163	84.680	31.936	5.515	1.00 10.00	A_13 A_13
ATOM		CD2 TYR	163	85.622	31.672	8.099	1.00 16.58	A_13
ATOM		CE2 TYR	163	84.266	31.867	7.873	1.00 12.32	A_13
ATOM		CZ TYR	163	83.807	31.991	6.576	1.00 11.77	A_13
ATOM		OH TYR	163	82.472	32.141	6.331	1.00 21.93	A_13
MOTA		C TYR	163	89.818	29.669	7.397	1.00 15.67	A_13
ATOM		O TYR	163	90.590	30.089	6.526	1.00 18.92	A_13
ATOM		N GLY	164	90.225	29.096	8.525	1.00 18.34	A_13
ATOM	1501	CA GLY	164	91.636	28.966	8.826	1.00 10.61	A_13
MOTA	1502	C GLY	164	92.149	30.215	9.525	1.00 15.63	A_13
MOTA	1503	O GLY	164	91.334	31.139	9.775	1.00 21.42	A_13
MOTA	1504	OT GLY	164	93.353	30.250	9.858	1.00 21.99	A_13
ATOM	3009	ZN ZN	166	73.275	35.223	18.371	1.00 27.40	AION
ATOM	3010	ZN ZN	167	65.511	41.122	10.564	1.00 27.86	AION
MOTA	3011 C		168	64.285	44.152	21.635	1.00 11.76	AION
ATOM		A CA	165	73.319	39.377	1.854	1.00 40.73	AION
ATOM	3017	C5 WAY	169	67.400	35.999	20.267	1.00 38.86	A693
MOTA	3018	CF1 WAY	169	66.626	35.606	19.161	1.00 30.96	A693
ATOM	3019	CH WAY	169	67.199	35.400	17.901	1.00 41.17	A693
MOTA	3020	C2 WAY	169	68.561	35.623	17.728	1.00 36.26	A693
ATOM	3021	C3 WAY	169	69.339	36.039	18.811	1.00 35.73	A693
MOTA MOTA	3022	C4 WAY N20 WAY	169	68.807	36.216	20.078	1.00 33.71	A693
MOTA	3023 3024	CD WAY	169 169	69.699	36.617	21.141	1.00 33.16	A693
ATOM	3024	C23 WAY	169	70.137	35.640	22.189	1.00 29.78	A693
MOTA	3025	C28 WAY	169	68.986 68.187	34.739 35.088	22.685 23.798	1.00 25.69 1.00 31.72	A693
MOTA	3027	C27 WAY	169	67.141	34.238	24.205	1.00 31.72	A693 A693
MOTA	3028	CM WAY	169	66.921	33.061	23.490	1.00 33.61	A693
ATOM	3029	N25 WAY	169	67.703	32.748	22.426	1.00 42.39	A693
ATOM	3030	C24 WAY	169	68.709	33.546	22.016	1.00 27.88	A693
ATOM	3031	S21 WAY	169	69.757	38.213	21.577	1.00 24.43	A693
ATOM	3032	C16 WAY	169	71.513	38.570	21.438	1.00 29.69	A693
ATOM	3033	C21 WAY	169	72.032	39.163	20.269	1.00 19.32	A693
MOTA	3034	C20 WAY	169	73.400	39.453	20.169	1.00 11.82	A693
MOTA	3035	C19 WAY	169	74.267	39.156	21.241	1.00 19.50	A693
MOTA	3036	C18 WAY	169	73.748	38.564	22.402	1.00 11.88	A693
MOTA	3037	C17 WAY	169	72.382	38.272	22.507	1.00 26.57	A693
MOTA	3038	O33 WAY	169	75.623	39.445	21.141	1.00 16.99	A693
ATOM	3039	C36 WAY	169	76.504	39.509	22.271	1.00 12.69	A693
MOTA	3040	O15 WAY	169	69.030	39.032	20.657	1.00 13.98	A693
MOTA	3041	014 WAY	169	69.419	38.338	22.942	1.00 22.94	A693
MOTA	3042	C7 WAY		70.780	36.256	18.621	1.00 30.48	A693
MOTA MOTA	3043 3044	N9 WAY	169 169	71.192	36.946 37.127	17.553	1.00 10.00	A693
MOTA	3045	YAW 80		72.581 71.614		17.426	1.00 38.25 1.00 39.46	A693
MOTA	3045	C29 WAY		66.584	35.847 36.175	19.414 21.566	1.00 39.46	A693 A693
ATOM	1505	CB THR		40.443	57.305	5.225	1.00 21.20	B_13
ATOM	1506	OG1 THR		39.149	56.999	-5.762	1.00 25.31	B_13
MOTA	1508	CG2 THR		41.017	56.087	4.541	1.00 23.15	B_13
ATOM	1509	C THR		40.920	59.113	6.901	1.00 32.45	B_13
MOTA	1510	O THR		41.453	59.582	7.908	1.00 36.97	B_13
ATOM	1513	N THR		41.386	56.786	7.488	1.00 34.12	B_13
MOTA	. 1515	CA THR		41.371	57.761	6.365	1.00 26.16	B_13
MOTA	1516	N LEU	8	39.907	59.694	6.265	1.00 23,60	B_13
MOTA	1518	CA LEU	8	39.387	60.984	6.649	1.00 22.66	B_13
MOTA	1519	CB LEU	8	38.113	60.848	7.503	1.00 21.78	B_13
MOTA	1520	CG LEU		36.860	61.484	6.863	1.00 27.13	·B_13
MOTA	1521	CD1 LEU	8	36.996	63.016	6.705	1.00 19.05	B_13
MOTA	1522	CD2 LEU		36.622	60.854	5.510	1.00 19.23	B_13
MOTA	1523	C LEU	8	40.432	61.896	7.298	1.00 27.16	B_13
MOTA	1524	O LEU	8	41.077	62.667	6.597	1.00 46.24	B_13
MOTA	1525	N LYS	9	40.615	61.804	8.618	1.00 27.84	B_13
ATOM	1527	CA LYS		41.572	62.674	9.306	1.00 15.20	B_13
MOTA	1528	CB LYS	, ,	41.147	64.143	9.148	1.00 32.32	B_13
MOTA MOTA	1529 1530	CG LYS		39.663	64.342	8.853	1.00 29.47	B_13
A1Un	1320	כה הוצ	7	38.788	64.243	10.084	1.00 28.34	B_13

ATOM	1531	CE	LYS	9	20 020	CC CCC	10 040	1 00 10 40	
					38.830	65.556	10.842	1.00 18.48	B_13
ATOM	1532	NZ	LYS	9	38.732	66.725	9.888	1.00 33.19	B_13
MOTA	1536	С	LYS	9	41.809	62.384	10.780	1.00 20.69	B_13
MOTA	1537	0	LYS	9	41.268	61.428	11.334	1.00 25.62	B_13
ATOM	1538	N	TRP	10	42.654	63.208	11.390	1.00 12.09	
ATOM	1540	CA	TRP						B_13
				10	42.988	63.112	12.813	1.00 21.78	B_13
MOTA	1541	CB	TRP	10	44.403	63.660	13.048	1.00 23.03	B_13
ATOM	1542	CG	TRP	10	45.499	62.890	12.349	1.00 27.60	B_13
ATOM	1543	CD2	TRP	10	46.077	61.650	12.762	1.00 27.28	B_13
ATOM	1544	CE2		10	47.071	61.302	11.829	1.00 22.11	B_13
MOTA	1545	CE3	TRP	10	45.859	60.781	13.847	1.00 11.66	B_13
ATOM	1546	CD1	TRP	10	46.153	63.247	11.198	1.00 21.84	B_13
MOTA	1547	NE1	TRP	10	47.094	62.305	10.873	1.00 10.00	B_13
ATOM	1549	CZ2							
				10	47.847	60.143	11.929	1.00 25.24	B_13
MOTA	1550	CZ3		10	46.632	59.622	13.951	1.00 22.71	B_13
ATOM	1551	CH2	TRP	10	47.611	59.317	12.999	1.00 15.23	B_13
ATOM	1552	C.	TRP	10	41.987	63.915	13.679	1.00 30.88	B_13
MOTA	1553	0	TRP	10	41.673	65.062	13.359	1.00 32.03	
									B_13
MOTA	1554	N	SER	11	41.495	63.316	14.765	1.00 35.64	B_13
ATOM	1556	CA	SER	11	40.548	63.981	15.665	1.00 30.37	B_13
ATOM	1557	CB	SER	11	39.498	62.995	16.176	1.00 31.03	B_13
ATOM	1558	OG	SER	11	38.485	62.815	15.202	1.00 41.11	B_13
ATOM	1560	c	SER	11.	41.206	64.691	16.840	1.00 20.70	5_13
									B_13
ATOM	1561	0	SER	11	40.558	65.002	17.838	1.00 36.52	B_13
MOTA	1562	N	LYS	12	42.504	64.910	16.731	1.00 23.56	B_13
ATOM	1564	CA	LYS	12	43.257	65.607	17.756	1.00 15.00	B_13
ATOM	1565	CB	LYS	12	43.991	64.631	18.688	1.00 18.58	B_13
ATOM	1566	CG	LYS	12	44.658	63.452	18.010		
								1.00 15.94	B_13
ATOM	1567	CD	LYS	12	45.456	62.589	.19.007	1.00 23.03	B_13
ATOM	1568	CE	LYS	12	44.593	61.715	19.933	1.00 27.10	B_13
ATOM	1569	NZ	LYS	12	44.075	62.402	21.157	1.00 34.75	B_13
MOTA	1573	С	LYS	12	44.200	66.453	16.914	1.00 25.03	B_13
MOTA	1574	ō	LYS	12	44.567				
						66.039	15.808	1.00 25.20	B_13
MOTA	1575	N	MET	13	44.536	67.647	17.401	1.00 18.44	B_13
MOTA	1577	CA	MET	13	45.377	68.582	16.663	1.00 24.63	B_13
ATOM	1578	CB	MET	13	44.864	70.015	16.880	1.00 13.15	B 13
ATOM	1579	CG	MET	13	43.421	70.253	16.419	1.00 21.56	
ATOM	1580								B_13
		SD	MET	13	43.167	70.131	14.616	1.00 31.39	B_13
MOTA	1581	CE	MET	13	41.433	69.678	14.474	1.00 24.70	B_13
MOTA	1582	С	MET	13	46.850	68.468	17.034	1.00 11.65	B_13
ATOM	1583	0	MET	13	47.728	68.815	16.247	1.00 14.33	B_13
ATOM	1584	N	ASN	14	47.103	67.985	18.242	1.00 16.99	B_13
ATOM	1586								
		CA	ASN	14	48.448	67.793	18.760	1.00 24.42	B_13
MOTA	1587	CB	ASN	14	48.437	68.006	20.268	1.00 17.84	B_13
ATOM	1588	CG	ASN	14	47.896	69.356	20.633	1.00 35.10	B_13
ATOM	1589	OD1	ASN	14	48.614	70.346	20.560	1.00 34.88	B_13
ATOM	1590		ASN	14	46.610	69.424	20.955	1.00 32.98	B_13
ATOM	1593		ASN	14					
		C			48.831	66.364	18.421	1.00 22.70	B_13
MOTA	1594	0	ASN	14	48.278	65.405	18.976	1.00 26.03	B_13
MOTA	1595	$\mathbf{N}$	LEU	15	49.706	66.228	17.432	1.00 18.07	B_13
MOTA	1597	CA	LEU	15	50.144	64.912	16.969	1.00 29.36	B_13
MOTA	1598	CB	LEU	15	49.878	64.775	15.466	1.00 24.35	B_13
ATOM	1599	CG	LEU	15	48.380		15.162		
ATOM	1600		LEU	15		64.762		1.00 19.51	B_13
					48.079	65.469	13.852	1.00 27.59	B_13
ATOM	1601		LEU	15	47.902	63.326	15.163	1.00 19.66	B_13
ATOM	1602	C	LEU	15	51.613	64.704	17.257	1.00 28.48	B_13
ATOM	1603	0	LEU	15	52.341	65.657	17.552	1.00 22.28	B_13
MOTA	1604	N	THR	16	52.044	63.453	17.198	1.00 12.77	B_13
MOTA	1606	CA	THR	16	53.433				
			1111			63.158	17.446	1.00 16.59	B_13
ATOM	1607	CB	THR	16	53.607	62.243	18.682	1.00 24.73	B_13
MOTA	1608		THR	16	52.912	61.005	18.481	1.00 12.79	B_13
MOTA	1610	CG2	THR	16	53.059	62.933	19.924	1.00 25.34	B_13
ATOM	1611	C	THR	16	54.038	62.515	16.214	1.00 21.94	B_13
ATOM	1612		THR						
		0		16	53.315	62.116	15.297	1.00 19.60	B_13
ATOM	1613	N	TYR	17	55.365	62.453	16.184	1.00 18.25	B_13
ATOM	1615	CA	TYR	17	56.092	61.810	15.097	1.00 19.54	B_13
ATOM	1616	CB	TYR	17	56.300	62.753	13.910	1.00 16.87	B_13
ATOM	1617	CG	TYR	17	57.277				
						63.892	14.116	1.00 27.90	B_13
MOTA	1618		TYR	17	56.839	65.135	14.587	1.00 13.93	B_13
ATOM	1619	CE1		17	57.700	66.221	14.652	1.00 17.08	B_13
ATOM	1620	CD2	TYR	17	58.613	63.764	13.723	1.00 14.99	B_13
ATOM	1621	CE2		17	59.479	64.841	13.777	1.00 25.98	B_13
ATOM	1622	CZ	TYR	17					
					59.017	66.075	14.242	1.00 33.12	B_13
MOTA	1623	OH	TYR	17	59.866	67.163	14.276	1.00 23.31	B_13
MOTA	1625	С	TYR	17	57.417	61.318	15.650	1.00 18.57	B_13
MOTA	1626	0	TYR	17	57.895	61.827	16.668	1.00 26.60	B_13
ATOM	1627	N	ARG	18	57.973	60.286	15.030	1.00 13.01	B_13

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ATOM	1629	CA	ARG	18	59.245	59.750	15.492	1.00 18.74	B_13
MOTA	1630	CB	ARG	18	59.033	58.589	16.473	1.00 11.96	B_13
MOTA	1631	CG	ARG	18	60.320	57.911	16.970	1.00 15.06	B_13
ATOM	1632	CD	ARG	18	60.012	56.596	17.690	1.00 11.72	B_13
MOTA	1633	NE	ARG	18	61.165	55.689	17.752	1.00 10.00	B_13
MOTA	1635	CZ	ARG	18	61.134	54.428	18.181	1.00 24.87	B_13
MOTA	1636	NH1	ARG	18	60.004	53.882	18.614	1.00 13.34	B_13
MOTA	1639	NH2	ARG	18	62.247	53.703	18.169	1.00 20.03	B_13
ATOM	1642	С	ARG	18	60.076	59.309	14.307	1.00 13.14	B_13
MOTA	1643	ō	ARG	18	59.598	58.588	13.434	1.00 14.10	
ATOM	1644	N	ILE	19	61.304	59.813			B_13
	1646						14.252	1.00 15.55	B_13
MOTA		CA	ILE	19	62.238	59.476	13.193	1.00 10.41	B_13
MOTA	1647	CB	ILE	19	63.307	60.603	13.054	1.00 17.20	B_13
MOTA	1648		ILE	19	64.273	60.307	11.903	1.00 16.57	B_13
MOTA	1649		ILE	19	62.613	61.952	12.836	1.00 15.47	B_13
MOTA	1650	CD1	ILE	19	63.543	63.110	12.783	1.00 14.99	B_13
MOTA	1651	C	ILE	19	62.870	58.166	13.673	1.00 10.00	B 13
MOTA	1652	0	ILE	19	63.829	58.179	14.434	1.00 10.00	B_13
ATOM	1653	N	VAL	20	62.289	57.037	13.276	1.00 17.84	B_13
ATOM	1655	CA	VAL	20	62.785	55.716	13.696	1.00 16.43	B 13
ATOM	1656	CB	VAL	20	61.911	54.570	13.138	1.00 13.17	_
ATOM	1657		VAL	20	62.519	53.208	13.493	1.00 10.00	B_13
MOTA	1658		VAL	20					B_13
MOTA					60.521	54.673	13.698	1.00 10.00	B_13
	1659	C	VAL	20	64.268	55.449	13.387	1.00 16.02	B_13
MOTA	1660	0	VAL	20	65.001	54.909	14.218	1.00 21.07	B_13
ATOM	1661	N	asn	21	64.698	55.762	12.177	1.00 10.00	B_13
MOTA	1663	CA	ASN	21	66.098	55.571	11.830	1.00 22.13	B_13
MOTA	1664	CB	ASN	21	66.392	54.128	11.386	1.00 19.75	B_13
MOTA	1665	CG	ASN	21	65.549	53.673	10.212	1.00 17.63	B_13
MOTA	1666	OD1	ASN	21	65.329	52.477	10.042	1.00 31.82	B_13
MOTA	1667	ND2	ASN	21	65.109	54.602	9.375	1.00 11.42	B_13
ATOM	1670	С	ASN	21	66.504	56.645	10.821	1.00 10.14	B_13
MOTA	1671	ŏ	ASN	21	65.639	57.377	10.340	1.00 11.74	
ATOM	1672	N	TYR	22	67.787	56.759	10.498		B_13
ATOM	1674	CA	TYR	22				1.00 12.25	B_13
ATOM	1675				68.233	57.829	9.602	1.00 12.46	B_13
		CB	TYR	22	69.136	58.800	10.383	1.00 23.15	B_13
ATOM	1676	CG	TYR	22	68.461	59.584	11.492	1.00 21.95	B_13
MOTA	1677		TYR	22	68.221	60.945	11.348	1.00 22.29	B_13
ATOM	1678	CE1		22	67.625	61.678	12.347	1.00 10.00	B_13
ATOM	1679	CD2	TYR	22	68.077	58.974	12.687	1.00 13.42	B_13
MOTA	1680	CE2	TYR	22	67.471	59.710	13.693	1.00 14.69	B_13
MOTA	1681	CZ	TYR	22	67.254	61.064	13.505	1.00 12.89	B_13
ATOM	1682	OH	TYR	22	66.660	61.829	14.466	1.00 16.56	B_13
ATOM	1684	С	TYR	22	68.988	57.395	8.359	1.00 11.62	B_13
ATOM	1685	0	TYR	22	69.793	56.478	8.407	1.00 16.23	B_13
MOTA	1686	N	THR	23	68.792	58.111	7.261	1.00 10.39	B_13
ATOM	1688	CA	THR	23	69.503	57.800		1.00 20.36	
MOTA	1689	CB	THR	23			6.024		B_13
MOTA	1690	OG1			68.909	58.582	4.829	1.00 16.21	B_13
ATOM				23	69.801	58.512	3.706	1.00 19.72	B_13
	1692	CG2		23	68.663	60.039	5.206	1.00 16.62	B_13
MOTA	1693	C	THR	23	70.990	58.153	6.163	1.00 17.35	B_13
MOTA	1694	0	THR	23	71.377	58.958	7.024	1.00 13.88	B_13
MOTA	1695	N	PRO	24	71.852	57.503	5.364	1.00 15.86	B_13
ATOM	1696	CD	PRO	24	71.625	56.247	4.629	1.00 17.29	B_13
MOTA	1697	CA	PRO	24	73.287	57.796	5.436	1.00 15.96	B_13
MOTA	1698	CB	PRO	24	73.920	56.570	4.763	1.00 10.00	B_13
ATOM	1699	CG	PRO	24	72.891	55.504	4.905	1.00 15.15	B_13
MOTA	1700	С	PRO	24	73.635	59.069	4.668	1.00 27.08	B_13
ATOM	1701	0	PRO	24	74.698	59.656	4.869	1.00 19.47	B_13
ATOM	1702	N	ASP	25	72.728	59.489	3.794	1.00 16.99	B_13
ATOM	1704	CA	ASP	25	72.927	60.663	2.958		
ATOM	1705	CB	ASP	25	71.792	60.758		1.00 10.00	B_13
ATOM	1706	CG					1.953	1.00 11.53	B_13
			ASP	25	71.665	59.521	1.105	1.00 33.88	B_13
MOTA	1707		ASP	25	70.570	59.311	0.556	1.00 22.66	B_13
MOTA	1708		ASP	25	72.653	58.762	0.980	1.00 29.59	B_13
ATOM	1709	C	ASP	25	73.068	62.011	3.642	1.00 23.36	B_13
MOTA	1710	0	ASP	25	73.694	62.916	3.093	1.00 20.32	B_13
ATOM	1711	N	MET	26	72.480	62.158	4.826	1.00 18.44	B_13
MOTA	1713	CA	MET	26	72.510	63.432	5.537	1.00 13.83	B_13
MOTA	1714	CB	MET	26	71.154	64.151	5.368	1.00 10.00	B_13
ATOM	1715	CG	MET	26	70.782	64.491	3.913	1.00 28.32	B_13
MOTA	1716	SD	MET	26	69.016	64.786	3.599	1.00 12.18	B_13
ATOM	1717	CE	MET	26	68.395	63.255	3.887	1.00 12.16	B_13
MOTA	1718	C	MET	26	72.827	63.238		1.00 37.25	
ATOM	1719	ŏ	MET	26			7.024		B_13
ATOM	1720	Ŋ	THR	27	72.839	62.107	7.533	1.00 20.90	B_13
ATOM					73.157	64.333	7.696	1.00 11.47	B_13
AT ON	1722	CA	THR	27	73.456	64.292	9.121	1.00 13.94	B_13

ATOM	1723	CB	THR	27	74.117	65.605	9.602	1.00 33.46	B_13
ATOM	1724	OGI	THR	27	73.209	66.702	9.415		
ATOM	1726		THR	27				1.00 10.00	B_13
					75.405	65.863	8.818	1.00 16.30	B_13
ATOM	1727	С	THR	27	72.135	64.113	9.861	1.00 10.67	B_13
ATOM	1728	0	THR	27	71.072	64.343	9.281	1.00 16.26	B_13
ATOM	1729	N	HIS	28	72.193	63.691	11.124	1.00 18.13	
ATOM	1731	CA	HIS	28					B_13
					70.986	63.514	11.915	1.00 10.00	B_13
ATOM	1732	CB	HIS	28	71.322	63.033	13.333	1.00 10.00	B_13
MOTA	1733	CG	HIS	28	71.793	61.608	13.401	1.00 22.65	B_13
ATOM	1734	CD2	HIS	28	72.893	61.003	12.889		
ATOM	1735							1.00 22.73	B_13
		ND1		28	71.103	60.627	14.080	1.00 19.90	B_13
MOTA	1737	CEl	HIS	28	71.755	59.481	13.985	1.00 16.52	B_13
ATOM	1738	NE2	HIS	28	72.843	59.681	13.268	1.00 20.38	B_13
ATOM	1740	С	HIS	28	70.281	64.870	11.957		
ATOM	1741							1.00 29.38	B_13
		0	HIS	28	69.074	64.941	11.742	1.00 17.20	B_13
ATOM	1742	N	SER	29	71.056	65.944	12.153	1.00 23.96	B_13
MOTA	1744	CA	SER	29	70.533	67.322	12.192	1.00 15.01	B_13
ATOM	1745	СВ	SER	29	71.661	68.334	12.438		
ATOM	1746	OG						1.00 14.05	B_13
			SER	29	72.117	68.303	13.770	1.00 18.32	B_13
MOTA	1748	С	SER	29	69.808	67.729	10.909	1.00 10.95	B_13
MOTA	1749	0	SER	29	68.732	68.314	10.971	1.00 24.24	B_13
ATOM	1750	N	GLU	30	70.415	67.449	9.757	1.00 10.96	B_13
ATOM	1752	CA	GLU	30	69.820	67.786			
ATOM	1753						8.470	1.00 10.00	B_:13
		CB	GLU	30	70.715	67.330	7.309	1.00 10.12	B_13
MOTA	1754	CG	GLU	30	71.967	68.143	7.042	1.00 22.31	B_13
ATOM	1755	CD	GLU	30	72.823	67.529	5.930	1.00 10.15	B_13
ATOM	1756	OE1	GLU	30	72.533	67.753	4.749		
ATOM	1757		GLU					1.00 31.98	B_13
				30	73.796	66.817	6.223	1.00 29.59	B_13
MOTA	1758	С	GLU	30	68.481	67.073	8.336	1.00 20.17	B_13
ATOM	1759	0	GLU	30	67.493	67.685	7.943	1.00 14.31	B 13
ATOM	1760	N	VAL	31	68.451	65.777	8.665	1.00 19.26	
ATOM	1762	CA	VAL						B_13
				31	67.228	64.989	8.536	1.00 14.22	B_13
MOTA	1763	CB	VAL	31	67.472	63.487	8.716	1.00 17.05	B_13
ATOM	1764	CG1	VAL	31	66.144	62.749	8.791	1.00 28.55	B_13
MOTA	1765	CG2	VAL	31	68.269	62.935	7.548	1.00 10.54	5_13
ATOM	1766	C	VAL	31					B_13
					66.138	65.458	9.477	1.00 12.36	B_13
ATOM	1767	0	VAL	31	64.963	65.488	9.093	1.00 12.83	B_13
ATOM	1768	N	GLU	32	66.530	65.805	10.703	1.00 20.46	B_13
ATOM	1770	CA	GLU	32	65.596	66.306	11.710	1.00 16.04	
MOTA	1771	CB	GLU	32					B_13
					66.269	66.365	13.094	1.00 14.71	B_13
MOTA	1772	CG	GLU	32	66.512	64.985	13.741	1.00 23.30	B_13
MOTA	1773	CD	GLŲ	32	67.724	64.930	14.700	1.00 21.41	B_13
ATOM	1774	OE1	GLU	32	68.229	63.823	15.003	1.00 15.79	
MOTA	1775	OE2	GLU	32					B_13
					68.183	65.985	15.157	1.00 13.71	B_13
MOTA	1776	C	GLU	32	65.125	67.697	11.257	1.00 27.19	B_13
MOTA	1777	0	GLU	32	63.951	68.042	11.383	1.00 19.82	B_13
MOTA	1778	N	LYS	33	66.021	68.461	10.636	1.00 12.52	B_13
MOTA	1780	CA	LYS	33	65.663	69.786	10.171		
ATOM	1781	CB	LYS					1.00 13.00	B_13
				33	66.889	70.592	9.762	1.00 22.63	B_13
MOTA	1782	CG	LYS	33	66.581	72.054	9.560	1.00 18.24	B_13
MOTA	1783	CD	LYS	33	65.604	72.545	10.630	1.00 29.21	B_13
ATOM	1784	CE	LYS	33	66.185	72.429	12.048	1.00 41.79	B_13
MOTA	1785	NZ	LYS	33	65.181	71.939			
MOTA	1789	c	LYS	33			13.054	1.00 20.17	B_13
					64.698	69.686	9.023	1.00 10.62	B_13
MOTA	1790	0	LYS	33	63.734	70.437	8.971	1.00 22.94	B_13
MOTA	1791	N	ALA	34	64.915	68.707	8.150	1.00 10.00	B_13
ATOM	1793	CA	ALA	34	64.050	68.475	7.000	1.00 11.94	B_13
MOTA	1794	CB	ALA	34	64.611	67.374			
ATOM	1795	c					6.100	1.00 10.00	B_13
			ALA	34	62.640	68.115	7.423	1.00 10.00	B_13
MOTA	1796	0	ALA	34	61.675	68.650	6.878	1.00 15.32	B_13
MOTA	1797	N	PHE	35	62.510	67.208	8.387	1.00 21.32	B_13
MOTA	1799	CA	PHE	35	61.187	66.789	8.852	1.00 18.32	D_13
MOTA	1800	CB	PHE						B_13
				35	61.267	65.451	9.614	1.00 25.48	B_13
ATOM	1801	CG	PHE	35	61.620	64.260	8.735	1.00 14.33	B_13
MOTA	1802	CD1	PHE	35	61.149	64.171	7.427	1.00 17.91	B_13
MOTA	1803	CD2	PHE	35	62.436	63.240	9.217	1.00 18.05	
MOTA	1804		PHE	35					B_13
					61.486	63.086	6.610	1.00 18.49	B_13
ATOM	1805		PHE	35	62.778	62.158	8.413	1.00 15.01	B_13
ATOM	1806	CZ	PHE	35	62.301	62.081	7.103	1.00 10.00	B_13
ATOM	1807	С	PHE	35	60.428	67.862	9.658	1.00 18.68	
ATOM	1808	ō	PHE	35			9.036		B_13
ATOM					59.202	67.971	9.556	1.00 17.05	B_13
	1809	N	LYS	36	61.160	68.664	10.425	1.00 16.30	B_13
MOTA	1811	CA	LYS	36	60.579	69.749	11.229	1.00 19.34	B_13
ATOM	1812	CB	LYS	36	61.676	70.420	12.052	1.00 24.61	B_13
ATOM	1813	CG	LYS	36	61.200	71.293	13.191	1.00 18.38	
ATOM	1814	CD	LYS	36					B_13
ATOM	1815				62.408	71.795	13.962	1.00 19.34	B_13
-31 011	T013	CE	LYS	36	62.067	72.267	15.356	1.00 21.80	B_13

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ATOM	1816	NZ	LYS	36	63.299	72.615	16.118	1.00 27.76	B_13
ATOM	1320	С	LYS	36	59.924	70.770	10.301		
								1.00 10.19	B_13
MOTA	1821	0	LYS	36	58.788	71.183	10.528	1.00 14.95	B_13
MOTA	1822	N	LYS	37	60.630	71.134	9.233	1.00 15.89	B_13
ATOM	1824	CA	LYS	37	60.126	72.076	8.230	1.00 19.95	
									B_13
ATOM	1825	CB	LYS	37	61.202	72.386	7.189	1.00 10.00	B_13
ATOM	1826	CG	LYS	37	62.209	73.439	7.569	1.00 13.18	B_13
MOTA	1827	CD	LYS	37	62.869	73.966	6.311	1.00 28.86	
									B_13
ATOM	1828	CE	LYS	37	61.825	74.460	5.281	1.00 31.44	B_13
MOTA	1829	NZ	LYS	37	60.878	75.512	5.772	1.00 26.23	B_13
MOTA	1833	C	LYS	37	58.939	71.482	7.472		
								1.00 25.64	B_13
ATOM	1834	0	LYS	37	57.968	72.177	7.161	1.00 24.39	B_13
ATOM	1835	N	ALA	38	59.060	70.205	7.128	1.00 17.12	B_13
MOTA	1837	CA	ALA	38	58.031	69.493			
		_					6.381	1.00 16.06	B_13
MOTA	1838	CB	ALA	38	58.459	68.038	6.154	1.00 12.19	B_13
ATOM	1839	С	ALA	38	56.692	69.557	7.094	1.00 11.12	B_13
ATOM	1840	0	ALA	38	55.648	69.736	6.458	1.00 31.10	
									B_13
MOTA	1841	N	PHE	39	56.732	69.393	8.417	1.00 21.01	B_13
MOTA	1843	CA	PHE	39	55.540	69.446	9.257	1.00 10.85	B_13
ATOM	1844	CB	PHE	39	55.841	68.833	10.639	1.00 14.45	
ATOM	1845								B_13
		:CG	PHE	39	55.851	67.325	10.659	1.00 21.88	B_13
ATOM	1846	CDI	PHE	39	57.016	66.625	10.954	1.00 16.88	B_13
MOTA	1847	CD2	PHE	39	54.675	66.599	10.442	1.00 22.14	B_13
ATOM	1848		PHE	39	57.010				
						65.223	11.037	1.00 17.95	B_13
ATOM	1849	CE2	PHE	39	54.655	65.190	10.522	1.00 17.22	B_13
MOTA	1850	CZ	PHE	39	55.823	64.503	10.823	1.00 13.51	B_13
ATOM.	1851	C	PHE	39	55.044	70.898	9.426		
								1.00 19.98	B_13
MOTA	1852	0	PHE	39	53.839	71.160	9.393	1.00 14.30	B_13
MOTA	1853	N	LYS	40	55.981	71.826	9.611	1.00 20.03	B_13
ATOM	1855	CA	LYS	40	55.681		9.795		
						73.245		1.00 18.64	B_13
ATOM	1856	CB	LYS	40	56.989	74.011	10.020	1.00 19.28	B_13
MOTA	1857	CG	LYS	40	57.064	75.392	9.440	1.00 26.34	B_13
MOTA	1858	CD	LYS	40	58.288	76.093	9.974	1.00 18.46	B_13
ATOM	1859								
		CE	LYS	40	58.021	76.673	11.339	1.00 20.86	B_13
MOTA	1860	NZ	LYS	40	57.053	77.814	11.232	1.00 27.28	B_13
MOTA	1864	С	LYS	40	54.899	73.790	8.612	1.00 20.57	B_13
ATOM	1865	Ō	LYS	40					
					54.034	74.654	8.756	1.00 22.54	B_13
MOTA	1866	N	VAL	41	55.216	73.251	7.445	1.00 17.15	B_13
ATOM	1868	ÇA	VAL	41	54.565	73.576	6.184	1.00 19.19	B_13
MOTA	1869	CB	VAL	41	55.095				
						72.566	5.086	1.00 17.28	B_13
MOTA	1870		VAL	41	53.987	72.064	4.160	1.00 10.00	B_13
ATOM	1871	CG2	VAL	41	56.224	73.191	4.293	1.00 19.38	· B_13
MOTA	1872	С	VAL	41	53.026	73.472	6.354		
ATOM								1.00 20.38	B_13
	1873	0	VAL	41	52.268	74.280	5.810	1.00 28.57	B_13
MOTA	1874	N	TRP	42	52.587	72.511	7.163	1.00 23.10	B_13
ATOM	1876	CA	TRP	42	51.166	72.265	7.403	1.00 19.29	B_13
ATOM	1877	CB	TRP	42					
					50.912	70.757	7.487	1.00 22.19	, B_13
MOTA	1878	CG	TRP	42	51.437	70.007	6.313	1.00 19.32	B_13
ATOM	1879	CD2	TRP	42	50.836	69.909	5.015	1.00 31.02	B_13
MOTA	1880	CE2	TRP	42	51.659	69.067	4.238	1.00 22.49	
ATOM	1881	CE3							
				42	49.677	70.448	4.434	1.00 15.54	B_13
MOTA	1882	CD1		42	52.571	69.251	6.269	1.00 14.04	B_13
MOTA	1883	NE1	TRP	42	52.710	68.681	5.027	1.00 13.55	B_13
ATOM	1885		TRP	42		. 68.752	2.912	1.00 18.87	
ATOM	1886				31.300				B_13
		C23	TRP	42	49.383	70.132	3.116	1.00 13.33	B_13
MOTA	1887	CH2	TRP	42	50.219	69.294	2.370	1.00 20.30	B_13
ATOM	1888	C	TRP	42	50.617	72.926	8.660	1.00 24.68	B_13
MOTA	1889	0	TRP	42	49.455	73.339			5_13
							8.688	1.0C 20.93	B_13
MOTA	1890	N	SER	43	51.432	72.987	9.710	1.00 20.63	B_13
MOTA	1892	CA	SER	43	51.007	73.601	10.968	1.00 22.47	B_13
MOTA	1893	CB	SER	43	51.955	73.231	12.116		
ATOM								1.00 10.00	B_13
	1894	OG	SER	43	53.265	73.716	11.891	1.00 33.50	B_13
MOTA	1896	С	SER	43	50.913	75.122	10.829	1.00 14.99	B_13
ATOM	1897	0	SER	43	50.224	75.784	11.595	1.00 11.58	5-13
ATOM	1898								B_13
		N	ASP	44	51.613	75.667	9.843	1.00 26.20	B_13
MOTA	1900	CA	ASP	44	51.595	77.100	9.617	1.00 22.11	B_13
ATOM	1901	CB	ASP	44	52.620	77.485	8.549	1.00 11.09	B_13
MOTA	1902	CG	ASP				0.343		
				44	54.000	77.751	9.125	1.00 18.45	B_13
ATOM	1903		ASP	44	54.903	78.114	8.347	1.00 17.67	B_13
MOTA	1904	OD2	ASP	44	54.195	77.602	10.345	1.00 21.44	B_13
MOTA	1905	С	ASP	44	50.216	77.575			5-13
ATOM	1906						9.190	1.00 32.83	B_13
		0	ASP	44	49.795	78.677	9.549	1.00 34.78	B_13
ATOM	1907	N	VAL	45	49.508	76.735	8.439	1.00 31.40	B_13
MOTA	1909	CA	VAL	45	48.191	77.094	7.932	1.00 14.00	B_13
ATOM	1910	СВ	VAL	45					5-13
					48.121	76.872	6.401	1.00 15.73	B_13
MOTA	1911		VAL	45	49.123	77.755	5.707	1.00 19.37	B_13
MOTA	1912	CG2	VAL	45	48.407	75.409	6.055	1.00 10.00	B_13
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ATOM	1913	С	VAL	45	47.054	76.333	8.575	1 00 10 41	
ATOM	1914							1.00 18.43	B_13
		0	VAL	45	45.954	76.304	8.026	1.00 26.09	B_13
ATOM	1915	N	THR	46	47.295	75.754	9.747	1.00 18.49	B_13
ATOM	1917	CA	THR	46	46.262	74.963	10.408		
								1.00 21.92	B_13
ATOM	1918	CB	THR	46	46.222	73.529	9.751	1.00 27.61	B_13
ATOM	1919	OG1	THR	46	44.876	73.047	9.661	1.00 28.78	B_13
ATOM	1921		THR						
				46	47.054	72.550	10.522	1.00 10.65	B_13
ATOM	1922	С	THR	46	46.505	74.931	11.932	1.00 18.41	B_13
ATOM	1923	0	THR	46	47.554				
						75.363	12.411	1.00 18.63	B_13
ATOM	1924	N	PRO	47	45.519	74.467	12.717	1.00 16.81	B_13
ATOM	1925	CD	PRO	47	44.113	74.209			
							12.348	1.00 32.80	B_13
MOTA	1926	CA	PRO	47	45.691	74.407	14.169	1.00 13.66	B_13
MOTA	1927	CB	PRO	47	44.256	74.489	14.675	1.00 30.52	B_13
MOTA	1928	CG	PRO	47					
					43.519	73.692	13.638	1.00 29.25	B_13
MOTA	1929	С	PRO	47	46.346	73.105	14.622	1.00 28.40	B_13
ATOM	1930	0	PRO	47	46.037	72.597	15.705		
								1.00 29.19	B_13
MOTA	1931	N	LEU	48	47.220	72.547	13.784	1.00 27.10	B_13
MOTA	1933	CA	LEU	48	47.915	71.302	14.124	1.00 21.49	B_13
MOTA	1934	CB	LEU	48					
					48.087	70.418	12.885	1.00 16.21	B_13
ATOM	1935	CG	LEU	48	46.924	69.476	12.538	1.00 15.14	B_13
MOTA	1936	CD1	LEU	48	45.618	70.049	13.000	1.00 26.83	
MOTA	1937								B_13
			LEU	48	46.894	69.206	11.035	1.00 32.93	B_13
ATOM	1938	С	LEU	48	49.262	71.611	14.771	1.00 16.35	B_13
ATOM	1939	0	LEU	48	49.885	72.648			5-13
							14.498	1.00 26.65	B_13
ATOM	1940	N	ASN	49	49.691	70.744	15.669	1.00 18.84	B_13
ATOM	1942	CA	ASN	49	50.956	70.940	16.354	1.00 25.67	5-13
ATOM	1943	СВ	ASN						B_13
				49	50.741	71.205	. 17.846	1.00 23.64	B_13
ATOM	1944	CG	ASN	49	49.734	72.301	18.100	1.00 23.64	B_13
MOTA	1945	001	ASN	49	48.895	72.192			
							18.989	1.00 33.47	B_13
MOTA	1946	MDZ	ASN	49	49.796	73.359	17.305	1.00 37.40	B_13
MOTA	1949	С	ASN	49	51.695	69.643	16.195	1.00 22.08	B_13
ATOM	1950	Ō	ASN						
				49	51.087	68.577	16.252	1.00 23.48	B_13
ATOM	1951	N	PHE	50	52.994	69.723	15.951	1.00 25.59	B_13
ATOM	1953	CA	PHE	50	53.762	68.510	15.806		
								1.00 19.57	B_13
ATOM	1954	СВ	PHE	50	54.258	68.343	14.380	1.00 12.47	B_13
MOTA	1955	CG	PHE	50	53.161	68.024	13.432	1.00 14.47	B_13
ATOM	1956		PHE						
				50	52.665	68.989	12.581	1.00 17.81	B_13
MOTA	1957	CD2	PHE	50	52.566	66.770	13.445	1.00 14.44	B_13
ATOM	1958	CE1	PHE	50	51.585	68.705			
							11.754	1.00 23.43	B_13
MOTA	1959	CEZ	PHE	50	51.488	66.482	12.624	1.00 20.62	B_13
MOTA	1960	CZ	PHE	50	50.999	67.447	11.781	1.00 13.34	
MOTA	1961								B_13
		C	PHE	50	.54.858	68.419	16.826	1.00 23.56	B_13
MOTA	1962	0	PHE	50	55.720	69.299	16.922	1.00 20.28	B_13
ATOM	1963	N	THR	51	54.728				
						67.387	17.651	1.00 26.45	B_13
MOTA	1965	CA	THR	51	55.650	67.090	18.725	1.00 29.37	B_13
MOTA	1966	CB	THR	51	54.851	66.834	20.024	1.00 28.17	B_13
ATOM	1967	OG1		51					
					53.946	65.738	19.824	1.00 40.86	B_13
ATOM	1969	CG2	THR	51	54.032	68.078	20.393	1.00 25.37	B_13
ATOM	1970	С	THR	51	56.435	65.838	18.331		
ATOM	1971							1.00 21.26	B_13
		0	THR	51	55.849	64.849	17.882	1.00 17.45	B_13
MOTA	1972	N	ARG	52	57.755	65.889	18.477	.1.00 15.17	B_13
ATOM	1974	CA	ARG	52	58.604				
						64.752	18.126	1.00 20.79	B_13
MOTA	1975	CB	ARG	52	59.868	65.241	17.429	1.00 20.81	B_13
ATOM	1976	CG	ARG	52	60.871	64.160	17.110	1.00 19.06	B_13
MOTA	1977	CD	ARG	52	62.208	64.808			
ATOM	1978						16.880	1.00 22.17	B_13
		NE	ARG	52	63.293	63.848	16.904	1.00 18.57	B_13
ATOM	1980	CZ	ARG	52	64.563	64.160	17.108	1.00 10.00	B_13
ATOM	1981	NHI	ARG	52	64.915	65.414	17.315		
								1.00 19.35	B_13
ATOM	1984		ARG	52	65.488	63.214	17.039	1.00 35.90	B_13
ATOM	1987	С	ARG	52	58.995	63.903	19.328	1.00 22.29	
MOTA	1988	ŏ	ARG	52					B_13
					59.326	64.433	20.387	1.00 24.98	B_13
MOTA	1989	N	LEU	53	59.013	62.586	19.140	1.00 19.90	B_13
ATOM	1991	CA	LEU	53	59.378				
						61.660	20.203	1.00 27.02	B_13
MOTA	1992	CB	LEU	53	58.279	60.625	20.434	1.00 16.80	B_13
MOTA	1993	CG	LEU	53	56.859	61.138	20.639	1.00 23.45	
ATOM	1994		LEU				20.033		B_13
				53	55.943	59.943	20.884	1.00 24.07	B_13
MOTA	1995	CD2	LEU	53	56.801	62.143	21.785	1.00 21.02	B_13
ATOM	1996	С	LEU	53	60.657				
ATOM						60.944	19.813	1.00 15.08	B_13
	1997	0	LEU	53	60.822	60.539	18.671	1.00 13.89	B_13
ATOM	1998	N	HIS	54	61.532	60.750	20.792	1.00 19.96	
MOTA	2000	CA	HIS	54					B_13
					62.812	60.079	20.568	1.00 28.80	B_13
ATOM	2001	CB	HIS	54	63.848	60.604	21.569	1.00 19.40	B_13
ATOM	2002	CG	HIS	54	64.113	62.075			
ATOM							21.431	1.00 31.96	B_13
	2003		HIS	54	63.365	63.060	20.883	1.00 21.32	B_13
ATOM	2004	ND1	HIS	54	65.292	62.662	21.835	1.00 33.94	B_13
ATOM	2006		HIS	,54	65.260				
						63.949	21.539	1.00 18.64	B_13
MOTA	2007	NE2	HIS	54	64.103	64.218	20.960	1.00 19.56	B_13
						•			

MOTA	2009	С	HIS	54	62.695	58.555	20.647	1.00 13.04	B_13
	2010		HIS	54					
ATOM					63.620	57.850	20.282	1.00 19.90	B_13
ATOM	2011		ASP	55	61.586	58.076	21.219	1.00 17.27	B_13
ATOM	2013	CA	ASP	55	61.303	56.648	21.366	1.00 25.79	B_13
ATOM	2014	CB	ASP	55	62.099	56.038	22.533	1.00 29.40	B 13
ATOM	2015	ĊĠ	ASP	55	63.443	55.428	22.076	1.00 29.64	_
									B_13
MOTA	2016	OD1	ASP	55	63.517	54.906	20.942	1.00 33.28	B_13
ATOM	2017	OD2	ASP	55	64.437	55.469	22.831	1.00 31.99	B_13
ATOM	2018	С	ASP	55	59.807	56.460	21.567	1.00 24.99	B_13
MOTA	2019	0	ASP	55	59.079	57.445	21.677	1.00 21.06	B_13
MOTA	2020	N	GLY	56	59.358	55.207	21.559	1.00 22.90	B_13
MOTA	2022	CA	GLY	56	57.954	54.877	21.737	1.00 21.80	B_13
ATOM	2023	C	GLY	56	57.155	54.926	20.447	1.00 14.48	
									B_13
MOTA	2024	0	GLY	56	57.720	55.108	19.379	1.00 19.38	B_13
ATOM	2025	N	ILE	57	55.841	54.742	20.545	1.00 11.78	B 13
ATOM	2027	CA	ILE	57	54.944	54.809	19.389	1.00 16.25	B_13
ATOM	202B	CB	ILE	57	53.737	53.804	19.510	1.00 22.94	B_13
MOTA	2029	CG2		57	52.442	54.417	18.955	1.00 24.79	B_13
MOTA	2030	CG1	ILE	57	54.025	52.505	18.744	1.00 25.63	B_13
MOTA	2031	CD1	TLE	57	53.586	52.520	17.240	1.00 17.48	B_13
ATOM	2032	c	ILE	57	54.410	56.238	19.301	1.00 18.78	
									B_13
MOTA	2033	0	ILE	57	53.866	56.777	20.270	1.00 11.40	B_13
ATOM	2034	N	ALA	58	54.598	56.842	18.140	1.00 14.67	B_13
ATOM	2036	CA	ALA	58	54.139	58.200	17.857	1.00 17.04	B_13
ATOM	2037	CB	ALA	58	55.270	59.015	17.245	1.00 10.00	B_13
MOTA	2038	С	ALA	58	53.048	58.009	16.825	1.00 25.41	B_13
ATOM	2039	0	ALA	58	52.956	56.940	16.243	1.00 22.59	B_13
MOTA	2040	N	ASP	59	52.211	59.020	16.609	1.00 13.36	B_13
ATOM	2042	CA	ASP	59	51.156	58.927	15.606	1.00 24.67	
									B_13
MOTA	2043	CB	ASP	59	50.348	60.237	15.545	1.00 10.00	B_13
ATOM	2044	CG	ASP	59	49.743	60.631	16.899	1.00 12.93	B_13
ATOM	2045	OD1	ASP	59	49.922	61.788	17.327	1.00 32.89	B_13
ATOM	2046		ASP	59	49.076	59.793	17.541	1.00 21.52	B_13
ATOM	2047	C	ASP	59	51.784	58.653	14.242	1.00 11.46	B_13
MOTA	2048	0	ASP	<b>59</b> .	51.378	57.736	13.531	1.00 16.58	B_13
MOTA	2049	N	ILE	60	52.791	59.445	13.899	1.00 24.90	B_13
ATOM	2051	CA	ILE	60	53.494	59.346			
							12.624	1.00 12.17	B_13
MOTA	2052	CB	ILE	60	53.620	60.738	11.975	1.00 10.91	B_13
ATOM	2053	CG2	ILE	60	54.289	60.641	10.588	1.00 10.70	B_13
ATOM	2054	CG1	ILE	60	52.228	61.367	11.851	1.00 18.58	B_13
ATOM	2055		ILE	60	52.219	62.870	11.726		
								1.00 12.00	B_13
MOTA	2056	С	ILE	60	54.881	58.750	12.841	1.00 12.93	B_13
ATOM	2057	0	ILE	60 ·	55.788	59.392	13.365	1.00 16.39	B_13
ATOM	2058	N	MET	61	55.015	57.485	12.483	1.00 19.08	B_13
ATOM	2060	CA	MET	61	56.275	56.784	12.617		B_13
								1.00 16.97	
MOTA	2061	CB	MET	61	56.011	55.328	13.035	1.00 23.79	B_13
ATOM	2062	CG	MET	61	55.313	55.172	14.422	1.00 12.37	B_13
ATOM	2063	SD	MET	61	56.389	55.360	15.913	1.00 31.01	B_13
ATOM	2064	CE	MET	61	57.204	53.749	15.861	1.00 14.93	B_13
MOTA									5_13
	2065	C	MET	61	56.995	56.888	11.265	1.00 12.72	B_13
ATOM	2066	0	MET	61	56.438	56.538	10.216	1.00 15.31	B_13
ATOM	2067	N	ILE	62	58.170	57.518	11.294	1.00 16.64	B_13
ATOM	2069	CA	ILE	62	58.978	57.739	10.097	1.00 27.48	B_13
MOTA	2070	CB	ILE.	62	59.557	59.181	10.060	1.00 10.00	B_13
						59.101			
MOTA	2071		ILE	62	60.191	59.462	8.717	1.00 18.65	B_13
MOTA	2072	CG1	ILE	62	58.460	60.203	10.342	1.00 18.51	B_13
MOTA	2073	CD1	ILE	62	58.983	61.499	10.931	1.00 16.23	B_13
MOTA	2074	С	ILE	62	60.155	56.787	10.046	1.00 15.06	B_13
ATOM									
	2075	0	ILE	62	60.873	56.606	11.033	1.00 10.73	B_13
MOTA	2076	N	SER	63	60.398	56.230	8.873	1.00 19.40	B_13
ATOM	2078	CA	SER	63	61.513	55.321	8.722	1.00 13.31	B_13
MOTA	2079	CB	SER	63	61.111	53.888	9.123	1.00 17.28	B_13
							9.123		D_13
MOTA	2080	OG	SER	63	59.985	53.435	8.391	1.00 13.66	B_13
MOTA	2082	С	SER	63	62.086	55.339	7.315	1.00 19.86	B_13
MOTA	2083	0	SER	63	61.441	55.766	6.347	1.00 20.93	B_13
MOTA	2084	N	PHE	64	63.338	54.914		1.00 17.78	
			Dir				7.237		B_13
MOTA	2086	CA	PHE	64	64.072	54.823	5.989	1.00 18.81	B_13
ATOM	2087	CB	PHE	64	65.409	55.553	6.105	1.00 16.50	B_13
MOTA	2088	CG	PHE	64	65.278	57.054	6.171	1.00 22.54	B_13
MOTA	2089		PHE						5-13
				64	65.321	57.817	5.013	1.00 20.48	B_13
ATOM	2090		PHE	64	65.155	57.708	7.395	1.00 24.76	B_13
ATOM	2091	CE1	PHE	64	65.246	59.207	5.071	1.00 13.94	B_13
ATOM	2092	CE2		64	65.079	59.105	7.461	1.00 14.29	B_13
ATOM	2093	CZ	PHE	64	65.128				
						59.847	6.298	1.00 10.16	B_13
MOTA	2094	C	PHE	64	64.293	53.336	5.823	1.00 10.30	B_13
MOTA	2095	0	PHE	64	64.571	52.637	6.799	1.00 14.11	B_13
MOTA	2096	N	GLY	65	64.121		4.610	1.00 13.58	B_13
		-	. – –	-					

MOTA	2098	CA	GLY	65	64 306	61 426	4 202		
					64.306	51.426	4.392	1.00 14.88	B_13
MOTA	2099	С	GLY	65	64.400	51.117	2.922	1.00 14.95	B_13
ATOM	2100	0	GLY	65	64.047	51.947	2.088	1.00 12.61	B_13
MOTA	2101	N	ILE	66	64.860	49.922	2.587	1.00 10.00	
ATOM	2103	CA	ILE	66	64.995	49.555			B_13
							1.187	1.00 19.70	B_13
ATOM	2104	CB	ILE	66	66.483	49.344	0.791	1.00 18.92	B_13
ATOM	2105	CG2	ILE	66	67.301	50.628	1.073	1.00 10.00	B_13
ATOM	2106	CG1	ILE	66	67.078	48.178	1.582	1.00 14.64	
ATOM	2107							1.00 14.04	B_13
			ILE	66	68.381	47.662	1.004	1.00 17.53	B_13
MOTA	2108	C	ILE	66	64.195	48.296	0.900	1.00 15.98	B_13
MOTA	2109	0	ILE	66	63.877	47.543	1.806	1.00 20.10	B_13
ATOM	2110	N	LYS	67	63.773	48.148			
							-0.349	1.00 18.78	B_13
MOTA	2112	CA	LYS	67	63.019	46.980	-0.787	1.00 14.73	B_13
MOTA	2113	СВ	LYS	67	63.986	45.827	-1.073	1.00 22.08	B_13
MOTA	2114	CG	LYS	67	65.107	46.142	-2.066	1.00 15.53	
MOTA	2115	CD	LYS	67					B_13
					64.591	46.325	-3.487	1.00 16.76	B_13
MOTA	2116	CE	LYS	67	65.573	45.763	-4.523	1.00 21.90	B_13
ATOM	2117	NZ	LYS	67	66.975	46.257	-4.394	1.00 28.03	B_13
MOTA	2121	С	LYS	67	61.945	46.548	0.218	1.00 16.24	
ATOM	2122		LYS	67					B_13
		0			61.136	47.360	0.649	1.00 10.25	B_13
MOTA	2123	N	GLU	68	61.968	45.293	0.630	1.00 10.00	B_13
ATOM	2125	CA	GLU	68	60.986	44.787	1.570	1.00 10.00	B_13
MOTA	2126	CB	GLU	68	61.004	43.257	1.505	1.00 31.44	B_13
ATOM	2127	CG	GLU	68					
					59.733	42.550	1.696	1.00 27.13	B_13
MOTA	2128	CD	GLU	68	58.723	42.720	0.524	1.00 12.88	B_13
ATOM	2129	OE1	GLU	68	59.106	42.180	-0.613	1.00 14.05	B_13
MOTA	2130		GLU	68	57.681	43.274	0.753		
ATOM	2131							1.00 38.61	B_13
		C	GLU	68	61.402	45.292	2.954	1.00 32.89	B_13
MOTA	2132	0	GLU	68	62.541	45.099	3.390	1.00 19.77	B_13
ATOM	2133	N	HIS	69	60.467	45.918	3.659	1.00 15.43	B_13
ATOM	2135	CA	HIS	69	60.777	46.473	4.964	1.00 10.00	
MOTA	2136	CB							B_13
			HIS	69	61.173	47.928	4.802	1.00 15.60	B_13
MOTA	2137	CG	HIS	69	60.151	48.731	4.063	1.00 18.06	B_13
MOTA	2138	CD2	HIS	69	59.131	49.509	4.498	1.00 25.01	B 13
ATOM	2139		HIS	69	60.055	48.709	2.689	1.00 21.79	
MOTA						40.703			B_13
	2141		HIS	69	59.023	49.430	2.308	1.00 19.43	B_13
MOTA	2142	NE2	HIS	69	58.438	49.932	3.384	1.00 19.23	B_13
MOTA	2143	С	HIS	69	59.655	46.396	5.978	1.00 16.27	B_13
ATOM	2144	0	HIS	69	59.689	47.099	6.969		
ATOM	2145							1.00 13.47	B_13
		N	GLY	70	58.610	45.629	5.719	1.00 21.21	B_13
ATOM	2147	CA	GLY	70	57.567	45.520	6.720	1.00 15.93	B_13
ATOM	2148	С	GLY	70	56.147	45.784	6.287	1.00 13.13	B_13
MOTA	2149	0	GLY	70	55.283	45.986	7.147		
ATOM	2150							1.00 12.19	B_13
		N	ASP	71	55.891	45.805	4.983	1.00 10.00	B_13
MOTA	2152	CA	ASP	71	54.540	46.030	4.480	1.00 17.84	B_13
MOTA	2153	CB	ASP	71	54.086	47.490	4.636	1.00 21.86	B_13
ATOM	2154	CG	ASP	71	54.946	48.480		1.00 13.38	
ATOM	2155		ASP				3.881		B_13
				71	54.896	49.644	4.291	1.00 10.00	B_13
MOTĄ	2156	OD2	ASP	71	55.633	48.135	2.897	1.00 10.00	B_13
ATOM	2157	С	ASP	71	54.313	45.557	3.064	1.00 27.18	B_13
ATOM	2158	0	ASP	71	55.221	45.068	2.416	1.00 16.61	
MOTA	2159	N	PHE	72	53.103				B_13
						45.759	2.564	1.00 10.00	B_13
MOTA	2161	CA	PHE	72	52.788	45.317	1.213	1.00 19.60	B_13
MOTA	2162	СB	PHE	72	51.292	45.017	1.099	1.00 16.43	B_13
MOTA	2163	CG	PHE	72	50.849	43.779	1.851	1.00 27.69	B_13
ATOM	2164		PHE	72	51.399				
ATOM						42.532	1.561	1.00 22.33	B_13
	2165		PHE	72	49.848	43.855	2.823	1.00 27.58	B_13
ATOM	2166		PHE	72	50.955	41.383	2.225	1.00 22.03	B_13
MOTA	2167	CE2	PHE	72	49.403	42.709	3.486	1.00 21.82	B_13
ATOM	2168	CZ	PHE	72	49.957		3.104		5_13
ATOM	2169					41.473	3.184	1.00 10.00	B_13
		C	PHE	72	53.225	46.313	0.130	1.00 18.56	B_13
MOTA	2170	0	PHE	72	52.840	46.190	-1.048	1.00 14.78	B_13
ATOM	2171	N	TYR	73	54.079	47.260	0.513	1.00 10.93	
ATOM	2173	CA	TYR	73	54.558				B_13
MOTA						48.295	-0.416	1.00 13.87	B_13
	2174	CB	TYR	73	53.943	49.649	-0.048	1.00 22.69	B_13
MOTA	2175	CG	TYR	73	52.439	49.581	0.007	1.00 16.43	B_13
MOTA	2176	CD1	TYR	73	51.774	49.385	1.219	1.00 18.21	B_13
ATOM	2177		TYR	73	50.386				
						49.219	1.257	1.00 35.13	B_13
MOTA	2178	CD2		73	51.683	49.618	-1.165	1.00 15.77	B_13
MOTA	2179	CE2	TYR	73	50.300	49.456	-1.133	1.00 39.16	B_13
ATOM	2180	CZ	TYR	73	49.663	49.258	0.080	1.00 28.27	B_13
ATOM	2181	ОН	TYR	73	48.301				
ATOM						49.122	0.106	1.00 33.06	B_13
	2183	C	TYR	73	56.088	48.349	-0.425	1.00 18.05	B_13
MOTA	2184	0	TYR	73	56.721	49.339	0.003	1.00 10.00	B_13
ATOM	2185	N	PRO	74	56.702	47.287	-0.953	1.00 13.76	B_13
MOTA	2186	CD	PRO	74	56.063				
ATOM						46.221	-1.740	1.00 14.21	B_13
and Old	2187	CA	PRO	. 74	58.158	47.183	-1.024	1.00 21.66	B_13

ATOM	2188	СВ	PRO	74	58.353	45.768	-1.569	1.00 15.88	B_13
MOTA	2189	CG	PRO	74	57.225	45.653	-2.540	1.00 13.95	B_13
ATOM	2190	C	PRO	74	58.747	48.226	-1.959	1.00 27.68	B_13
ATOM ATOM	2191 2192	N O	PRO PHE	74 75	58.173 59.883	48.526 48.794	-3.012 -1.562	1.00 21.90 1.00 20.91	B_13 B_13
ATOM	2194	CA	PHE	75	60.554	49.773	-2.395	1.00 15.84	B_13
MOTA	2195	CB	PHE	75	61.498	50.637	-1.548	1.00 11.67	B_13
MOTA	2196	CG.	PHE	75	60.765	51.589	-0.641	1.00 14.42	B_13
ATOM ATOM	2197 2198		PHE PHE	75 75	59.831	52.484	-1.162 0.726	1.00 16.56	B_13
ATOM	2199		PHE	75 75	60.976 59.119	51.574 53.345	-0.327	1.00 10.00 1.00 11.14	B_13 B_13
MOTA	2200		PHE	75	60.274	52.423	1.558	1.00 10.28	B_13
ATOM	2201	CZ	PHE	75	59.340	53.316	1.027	1.00 10.00	B_13
MOTA	2202	C	PHE	75 75	61.236	49.068	-3.573	1.00 14.23	B_13
ATOM ATOM	2203 2204	O N	PHE ASP	75 76	61.357 61.742	47.837 49.845	-3.582 -4.526	1.00 18.64 1.00 12.83	B_13 B_13
ATOM	2206	CA	ASP	76	62.330	49.287	-5.740	1.00 20.69	B_13
MOTA	2207	CB	ASP	76	61.394	49.644	-6.911	1.00 14.28	B_13
MOTA	2208	CG	ASP	76 76	61.212	51.144	-7.080	1.00 14.37	B_13
ATOM ATOM	2209 2210		ASP ASP	76 76	61.361 60.941	51.882 51.597	-6.095 -8.202	1.00 22.32 1.00 15.92	B_13 B_13
ATOM	2211	C	ASP	76	63.764	49.698	-6.104	1.00 19.31	B_13
ATOM	2212	Ó	ASP	76	64.056	49.864	-7.278	1.00 18.67	B_13
ATOM	2213	N	GLY	77	64.653	49.902	-5.132	1.00 10.00	B_13
MOTA MOTA	2215 2216	CA C	GLY GLY	77 77	65.997	50.326	-5.501	1.00 10.00	B_13
ATOM	2217	ŏ	GLY	77	65.989 64.967	51.790 52.487	-5.970 -5.752	1.00 16.22 1.00 17.04	B_13 B_13
MOTA	2218	Ň	PRO	78	67.080	52.305	-6.589	1.00 12.53	B_13
MOTA	2219	CD	PRO	78	68.319	51.564	-6.856	1.00 12.24	B_13
MOTA	2220	CA	PRO	78 70	67.207	53.691	-7.086	1.00 11.81	B_13
MOTA MOTA	2221 2222	CB CG	PRO PRO	78 78	68.546 69.316	53.678 52.693	-7.816 -7.066	1.00 10.00 1.00 12.78	B_13 B_13
ATOM	2223	c	PRO	78	66.093	54.146	-8.027	1.00 10.00	B_13
MOTA	2224	0	PRO	78	65.621	53.381	-8.853	1.00 27.46	B_13
MOTA	2225	N	SER	79	65.641	55.386	-7.852	1.00 19.14	B_13
MOTA MOTA	2227 2228	CA	SER	79 79	64.568	55.963	-8.669	1.00 10.00	B_13
ATOM	2229	CB OG	SER SER	79 79	64.970 63.982		-10.148 -10.901	1.00 20.11 1.00 23.87	B_13 B_13
ATOM	2231	C	SER	79	63.231		-8.507	1.00 31.68	B_13
ATOM	2232	0	SER	79	63.074	54.356	-7.606	1.00 26.48	B_13
ATOM	2233	N	GLY	80	62.250	55.589	-9.327	1.00 10.00	B_13
ATOM ATOM	2235 2236	CA C	GLY	80 80	60.940 60.293	54.969 55.412	-9.260 -7.968	1.00 10.07 1.00 30.72	B_13
ATOM	2237	ŏ	GLY	80	60.347	56.600	-7.643	1.00 30.72	B_13 B_13
MOTA	2238	N	LEU	81	59.779	54.452	-7.193	1.00 23.74	B_13
ATOM	2240	CA	LEU	81	59.135	54.752	-5.917	1.00 13.14	B_13
MOTA MOTA	2241 2242	CB CG	LEU	81 81	58.661	53.481	-5.213	1.00 16.20	B_13
ATOM	2243		LEU	81	57.393 57.554	52.775 52.277	-5.687 -7.096	1.00 17.33 1.00 28.67	B_13 B_13
ATOM	2244		LEU	81	57.103	51.617	-4.745	1.00 27.02	B_13
ATOM	2245	C	LEU	81	60.122	55.466		1.00 14.51	B_13
MOTA MOTA	2246 2247	N	LEU	81 82	61.264	55.016	-4.846	1.00 16.24	B_13
ATOM	2249	CA	LEU	82	59.692 60.540	56.590 57.381	-4.470 -3.594	1.00 11.33 1.00 17.52	B_13 B_13
MOTA	2250	CB	LEU	82	60.442	58.861	-3.986	1.00 18.51	B_13
ATOM	2251	CG	LEU	82	61.355	59.499	-5.044	1.00 15.37	B_13
MOTA MOTA	2252 2253		LEU	82 82	61.800 60.639	58.504		1.00 17.05	B_13
MOTA	2254	C	LEU	82	60.172	60.744 57.203	-5.659 -2.127	1.00 16.87 1.00 10.00	B_13 . B_13
MOTA	2255	O	LEU	82	61.045	57.056		1.00 19.90	B_13
MOTA	2256	N	ALA	83	58.876	57.201	-1.840	1.00 18.16	B_13
ATOM ATOM	2258 2259	CA	ALA	83	58.378	57.077		1.00 13.17	B_13
ATOM	2259	CB C	ALA ALA	83 83	58.762 56.846	58.322 56.925		1.00 10.00	B_13 B_13
MOTA	2261	ō	ALA	83	56.209	57.155		1.00 10.00	B_13
MOTA	2262	N	HIS	84	56.268	56.619		1.00 10.00	B_13
ATOM	2264	CA	HIS	84	54.811	56.472		1.00 23.81	B_13
ATOM ATOM	2265	CB	HIS	84	54.270	55.188		1.00 30.45	B_13
MOTA	2266 2267	CG	HIS HIS	84 84	54.848 54.856	53.925 53.415		1.00 17.68	B_13 B_13
ATOM	2268		HIS	84	55.525	53.025	-0.076	1.00 10.00	B_13
MOTA	2270	CE:	l HIS	84	55.933	52.015	0.666	1.00 29.72	B_13
MOTA	2271		HIS	84	55.543	52.224		1.00 13.81	B_13
MOTA MOTA	2272 2273	C	HIS HIS	84 84	54.363 55.099	56.547 56.148		1.00 12.82	B_13
MOTA	2274	N	ALA	85	53.161	57.076		1.00 20.02 1.00 28.38	B_13 B_13
ATOM	2276	CA	ALA	85	52.584	57.230		1.00 18.64	B_13

ATOM	2277	СВ	ALA	85	52.638	58.705	4.223	1.00 13.89	B_13
ATOM	2278	C	ALA	85	51.138	56.716	3.837	1.00 10.00	
				85	50.434				B_13
ATOM	2279	0	ALA			56.728	2.828	1.00 10.00	B_13
ATOM	2280	N	PHE	86	50.676	56.322	5.016	1.00 14.76	B_13
ATOM	2282	CA	PHE	86	49.316	55,811	5.143	1.00 17.96	B_13
ATOM	2283	CB	PHE	86	49.286	54.592	6.084	1.00 15.86	B_13
ATOM	2284	CG	PHE	86	50.320	53.542	5.748	1.00 26.30	
									B_13
ATOM	2285		PHE	86	49.973	52.398	5.042	1.00 22.30	B_13
ATOM	2286	CD2	PHE	86	. 51 . 654	53.730	6.090	1.00 27.63	B_13
ATOM	2287	CE1	PHE	86	50.938	51.472	4.681	1.00 27.85	B_13
ATOM	2288	CE2	PHE	86					
					52.620	52.810	5.731	1.00 13.97	B_13
ATOM .	2289	CZ	PHE	86	52.266	51.683	5.027	1.00 23.08	B_13
ATOM	2290	С	PHE	86	48.427	56.924	5.669	1.00 13.02	B_13
ATOM	2291	0	PHE	86	48.870	57.747	6.466	1.00 15.02	B 13
MOTA	2292	N	PRO	87	47.174	57.006	5.186	1.00 17.55	
									B_13
MOTA	2293	CD	PRO	87	46.565	56.165	4.146	1.00 10.17	B_13
ATOM	2294	CA	PRO	87	46.228	58.041	5.628	1.00 32.09	B_13
ATOM	2295	CB	PRO	87	44.961	57.720	4.819	1.00 18.55	B_13
MOTA	2296	CG	PRO	87	45.115	56.277	4.481	1.00 18.86	B_13
ATOM	2297	c	PRO	87		57.955	7.139		
	-				45.995			1.00 25.18	B_13
MOTA	2298	0	PRO	87	46.284	56.919	7.752	1.00 18.18	B_13
ATOM	2299	N	PRO	88	45.462	59.032	7.760	1.00 11.49	B_13
ATOM	2300	CD	PRO	88	45.015	60.303	7.164	1.00 10.00	B_13
ATOM	2301	CA	PRO	88	45.217	59.034	9.202	1.00 19.03	B_13
ATOM	2302		PRO	88		60.302			
		CB			44.399		9.402	1.00 14.16	B_13
MOTA	2303	CG	PRO	88	44.939	61.196	8.357	1.00 16.39	B_13
MOTA	2304	С	PRO	88	44.500	57.787	9.733	1.00 25.43	B_13
MOTA	2305	0	PRO	88	43.670	57.165	9.044	1.00 15.90	B_13
ATOM	2306	N	GLY	89	44.865	57.422	10.955	1.00 26.28	
									B_13
MOTA	2308	CA	GLY	89	44.299	56.264	11.606	1.00 25.32	B_13
ATOM	2309	C	GLY	89	45.343	55.713	12.546	1.00 34.38	B_13
ATOM	2310	0	GLY	89	46.485	56.164	12.498	1.00 23.28	B_13
ATOM	2311	N	PRO	90	44.977	54.774	13.437	1.00 13.87	
MOTA	2312	CD							B_13
			PRO	90	43.613	54.259	13.631	1.00 16.36	B_13
MOTA	2313	CA	PRO	90	45.898	54.164	14.398	1.00 10.34	B_13
MOTA	2314 ·	CB	PRO	90	44.963	53.360	15.300	1.00 15.93	B_13
ATOM	. 2315	CG	PRO	90	43.870	52.975	14.373	1.00 23.25	B_13
ATOM	2316								
		C	PRO	90	46.942	53.299	13.711	1.00 18.38	B_13
MOTA	2317	0	PRO	90	46.875	53.064	12.505	1.00 26.81	B_13
MOTA	2318	N	ASN	91	47.903	52.831	14.502	1.00 26.63	B_13
MOTA	2320	CA	ASN	91	49.022	52.010	14.033	1.00 21.91	B_13
ATOM	2321	CB	ASN	91	48.740	50.500	14.081	1.00 18.89	B_13
ATOM	2322	CG	ASN	91	47.437	50.117	13.448	1.00 22.49	B_13
MOTA	2323		ASN	91	47.335	50.017	12.237	1.00 29.37	B_13
ATOM	2324	ND2	ASN	91	46.438	49.858	14.273	1.00 28.01	B_13
MOTA	2327	С	ASN	91	49.656	52.438	12.721	1.00 20.07	B_13
ATOM	2328	0	ASN	91	50.301	53.479	12.681	1.00 21.24	B_13
ATOM	2329	N	TYR	92					
					49.423	51.716	11.633	1.00 20.15	B_13
ATOM	2331	CA	TYR	92	50.052	52.081	10.367	1.00 18.70	B_13
ATOM	2332	CB	TYR	92	49.905	50.953	9.344	1.00 14.48	B_13
MOTA	2333	CG	TYR	92	50.906	49.821	9.567	1.00 24.41	B_13
ATOM	2334	CD1	TYR	92	52.266	50.003	9.287	1.00 27.39	B_13
MOTA	2335		TYR	92	53.198	48.979	9.471	1.00 18.14	
									B_13
ATOM	2336	CD2		92	50.499	48.571	10.044	1.00 28.07	B_13
MOTA	2337	CE2		92	51.427	47.529	10.230	1.00 36.50	B_13
MOTA	2338	cz	TYR	92	52.778	47.741	9.940	1.00 43.64	B_13
ATOM	2339	OH	TYR	92	53.694	46.710	10.105	1.00 32.21	B_13
MOTA	2341	С	TYR	92	49.633	53.431	9.797	1.00 21.78	B_13
ATOM	2342	ŏ	TYR	92					
					50.384	54.049	9.040	1.00 12.55	B_13
MOTA	2343	N	GLY	93	48.464	53.916	10.198	1.00 15.83	B_13
MOTA	2345	CA	GLY	93	48.015	55.216	9.732	1.00 11.69	B_13
ATOM	2346	С	GLY	93	48.971	56.326	10.134	1.00 18.60	B_13
ATOM	2347	0	GLY	93	49.561	56.300	11.227	1.00 22.00	
ATOM	2348		GLY	94					B_13
		N			49.205	57.258	9.216	1.00 10.27	B_13
MOTA	2350	CA	GLY	94	50.099	58.365	9.492	1.00 18.36	B_13
ATOM	2351	C	GLY	94	51.567	58.061	9.234	1.00 15.54	· B_13
MOTA	2352	Ó	GLY	94	52.334	58.967	8.938	1.00 17.55	B_13
ATOM	2353	N	ASP	95	51.977	56.801	0.250	1.00 17.69	P_T3
							9.351		B_13
ATOM	2355	CA	ASP	95	53.386	56.457	9.134	1.00 19.67	B_13
ATOM	2356	CB	ASP	95	53.637	54.986	9.444	1.00 15.96	B_13
MOTA	2357	CG	ASP	95	53.346	54.634	10.900	1.00 25.37	B_13
ATOM	2358		ASP	95	53.627	53.484	11.297	1.00 16.05	B_13
ATOM	2359		ASP	95					
					52.835	55.488	11.656	1.00 14.66	B_13
ATOM	2360	C	ASP	95	53.896	56.808	7.733	1.00 17.15	B_13
ATOM	2361	0	ASP	95	53.162	56.711	6.746	1.00 19.09	B_13
ATOM	2362	N	ALA	96	55.166	57.198	7.662	1.00 18.71	B_13
ATOM	2364	CA	ALA	96	55.803	57.581	6.400	1.00 19.97	B_13
				-			3.100	,,	~

ATOM	2365	СВ	ALA	96	56.098	59.095	6.379	1.00 22.61	D 13
ATOM	2366		ALA	96	57.088	56.784	6.204	1.00 25.63	B_13 B_13
ATOM	2367		ALA	96	57.948		. 7.095	1.00 12.54	B_13
ATOM	2368		HIS	97	57.211	56.166	5.035	1.00 13.27	B_13
ATOM	2370		HIS	97	58.375	55.357	4.730	1.00 25.28	B_13
ATOM	2371		HIS	97	57.955	53.905	4.464	1.00 10.00	B_13
ATOM	2372		HIS	97	57.264	53.257	5.624	1.00 12.02	B_13
ATOM	2373	CD2		97	57.214	53,603	6.929	1.00 10.00	B_13
ATOM	2374	ND1		97	56.516	52.104	5.499	1.00 12.91	B_13
ATOM	2375	CE1		97	56.038	51.770	6.688	1.00 10.00	B_13
ATOM	2376	NE2		97	56.445	52.664	7.571	1.00 10.64	B_13
ATOM	2378	C	HIS	97	59.069	55.959	3.520	1.00 13.82	B_13
ATOM	2379	ŏ	HIS	97	58.415	56.273	2.517	1.00 12.27	B_13
ATOM	2380	N	PHE	98	60.379	56.154	3.647	1.00 10.67	B_13
MOTA	2382	CA	PHE	98	61.224	56.718	2.595	1.00 15.67	B_13
ATOM	2383	СВ	PHE	98	61.970	57.938	3.156	1.00 10.76	B_13
ATOM	2384	CG	PHE	98	61.055	59.025	3.627	1.00 17.93	B_13
ATOM	2385	CD1		98	60.730	60.082	2.786	1.00 17.93	B_13
ATOM	2386	CD2		98	60.476	58.974	4.893	1.00 14.14	B_13
ATOM	2387		PHE	98	59.833	61.066	3.201	1.00 22.42	B_13
ATOM	2388	CE2	PHE	98	59.574	59.962	5.315	1.00 10.00	B_13
ATOM	2389	CZ	PHE	98	59.257	61.002	4.469	1.00 10.00	B_13
ATOM	2390	c	PHE	98	62.218	55.669	2.064	1.00 26.64	B_13
ATOM	2391	ō	PHE	98	62.882	54.969	2.851	1.00 13.27	B_13
ATOM	2392	N	ASP	99	62.331	55.577	0.738	1.00 12.24	B_13
ATOM	2394	CA	ASP	99	63.229	54.612	0.102	1.00 10.00	B_13
ATOM	2395	СВ	ASP	99	62.884	54.471	-1.385	1.00 10.00	B_13
ATOM	2396	CG	ASP	99	63.615	53.311	-2.067	1.00 22.86	B_13
MOTA	2397	OD1		99	63.170	52.890	-3.160	1.00 11.60	B_13
ATOM	2398		ASP	99	64.624	52.806	-1.528	1.00 21.20	B_13
MOTA	2399	С	ASP	99	64.677	55.046	0.264	1.00 12.66	B_13
MOTA	2400	0	ASP	99	65.121	56.010	-0.366	1.00 18.37	B_13
ATOM	2401	N	ASP	100	65.439	54.289	1.046	1.00 12.86	B_13
MOTA	2403	CA	ASP	100	66.833	54.642	1.260	1.00 14.46	B_13
MOTA	2404	CB	ASP	100	67.308	54.271	2.660	1.00 17.70	B_13
ATOM	2405	CG	ASP	100	68.006	55.437	3.358	1.00 16.15	B_13
ATOM	2406		ASP	100	68.091	55.447	4.602	1.00 15.74	B_13
MOTA	2407		ASP	100	68.470	56.354	2.655	1.00 27.08	B_13
ATOM	2408	c	ASP	100	67.793	54.171	0.179	1.00 13.66	B_13
ATOM	2409	ō	ASP	100	68.961	53.932	0.416	1.00 19.54	B_13
MOTA	2410	N	ASP	101	67.254	53.954	-1.010	1.00 12.83	B_13
MOTA	2412	CA	ASP	101	68.074	53.590	-2.164	1.00 10.00	B_13
ATOM	2413	CB	ASP	101	67.471	52.413	-2.933	1.00 10.00	B_13
ATOM	2414	CG	ASP	101	67.997	51.065	-2.449	1.00 16.87	B_13
ATOM	2415		ASP	101	67.232	50.089	-2.458	1.00 19.89	B_13
ATOM	2416		ASP	101	69.184	50.968	-2.066	1.00 18.51	B_13
ATOM	2417	C	ASP	101	68.108	54.858	-3.029	1.00 26.72	B_13
ATOM	2418	Õ	ASP	101	68.602	54.853	-4.172	1.00 12.11	B_13
ATOM	2419	N	GLU	102	67.500	55.922	-2.496	1.00 13.76	B_13
ATOM	2421	CA	GLU	102	67.462	57.217	-3.161	1.00 12.54	B_13
ATOM	2422	CB	GLU	102	66.135	57.958	-2.916	1.00 13.01	B_13
ATOM	2423	CG	GLU	102	64.873	57.257	-3.381	1.00 15.50	B_13
ATOM	2424	CD	GLU	102	64.973	56.707	-4.791		B 13
ATOM	2425		GLU	102	65.640	57.307	-5.665	1.00 12.78	B_13
MOTA	2426		GLU	102	64.399	55.635	-5.021	1.00 12.36	B_13
ATOM	2427	С	GLU	102	68.544	58.040	-2.505	1.00 14.96	B_13
MOTA	2428	0	GLU	102	68.939	57.760	-1.371	1.00 10.00	B_13
ATOM	2429	N	THR	103	69.030	59.039	-3.228	1.00 19.38	B_13
ATOM	2431	CA	THR	103	70.021	59.957	-2.693	1.00 16.49	B_13
ATOM	2432	CB	THR	103	70.973	60.490	-3.801	1.00 19.31	B_13
ATOM	2433	OG1	THR	103	71.661	59.384	-4.399	1.00 25.44	B_13
ATOM	2435	CG2	THR	103	72.006	61.462	-3.212	1.00 10.75	B_13
MOTA	2436	С	THR	103	69.180	61.104	-2.141	1.00 12.91	B_13
MOTA	2437	0	THR	103	68.414	61.727	-2.867	1.00 13.59	B_13
MOTA	2438	N	TRP	104	69.252	61.322	-0.842	1.00 20.60	B_13
MOTA	2440	CA	TRP	104	68.497	62.388	-0.237	1.00 20.00	B_13
MOTA	2441	CB	TRP	104	67.852	61.902	1.063	1.00 22.66	B_13
ATOM	2442	CG	TRP	104	66.837	60.808	0.870	1.00 22.00	B_13
ATOM	2443		TRP	104	65.505	60.953	0.347	1.00 27.35	B_13
ATOM	2444		TRP	104	64.936	59.654	0.287	1.00 27.33	B_13 B_13
ATOM	2445	CE3		104	64.741	62.054	-0.079	1.00 12.81	B_13
ATOM	2446		TRP	104	67.013	59.473	1.108	1.00 17.89	B_13
ATOM	2447		TRP	104	65.876	58.775	0.755	1.00 17.89	B_13
ATOM	2449		TRP	104	63.632	59.429	-0.186	1.00 10.00	B_13
MOTA	2450		TRP	104	63.445	61.832	-0.549	1.00 22.21	B_13
ATOM	2451		TRP	104	62.904	60.527	-0.598	1.00 23.31	B_13
MOTA	2452	c	TRP	104	69.416	63.570	0.033	1.00 16.43	B_13
		-				55.570	3.033	2.00 20.43	<i></i> 13

MOTA	2453	0	TRP	104	70.520	63.380	0.526	1.00 11.13	D 13
	_								B_13
ATOM	2454		THR	105	68.960	64.775	-0.322	1.00 19.48	B_13
ATOM	2456	CA	THR	105	69.716	66.015	-0.097	1.00 10.40	B_13
MOTA	2457	CB	THR	105	70.153	66.749	-1.398	1.00 10.00	B_13
ATOM	2458	OG1	THR	105	69.305	66.401	-2.501	1.00 18.53	B_13
ATOM	2460		THR	105	71.596	66.484	-1.709	1.00 34.62	
									B_13
ATOM	2461	С	THR	105	68.904	67.062	0.641	1.00 20.82	B_13
MOTA	2462	0	THR	105	67.686	66.952	0.768	1.00 15.93	B_13
MOTA	2463	N	SER	106	69.621	68.073	1.125	1.00 38.37	B_13
ATOM	2465	CA	SER	106	69.029	69.222	1.791	1.00 20.77	B_13
MOTA	2466	CB	SER	106	69.979	69.778	2.862	1.00 17.95	B_13
MOTA	2467	OG	SER	106	70.281	68.825	3.864	1.00 29.88	B_13
ATOM	2469	C	SER	106	68.889	70.245	0.657	1.00 19.23	B_13
	2470								
MOTA		0	SER	106	68.202	71.260	0.782	1.00 21.34	B_13
MOTA	2471	N	SER	107	69.577	69.981	-0.450	1.00 18.73	B_13
ATOM	2473	CA	SER	107	69.533	70.884	-1.592	1.00 20.92	B_13
MOTA	2474	CB	SER	107	70.945	71.380	-1.927	1.00 19.84	B_13
	2475								
MOTA		OG	SER	107	71.556	71.957	-0.788	1.00 27.31	B_13
MOTA	2477	C	SER	107	68.848	70.284	-2.828	1.00 18.68	B_13
ATOM	2478	0	SER	107	67.660	69.953	-2.771	1.00 21.51	B_13
ATOM	2479	N	SER	108	69.623	70.038	-3.888	1.00 18.53	B_13
	2481		SER						5-13
ATOM		CA		108	69.091	69.544	-5.152	1.00 16.21	B_13
MOTA	2482	CB	SER	108	69.285	70.632	-6.205	1.00 29.10	B_13
ATOM	2483	OG	SER	108	70.665	70.969	-6.271	1.00 21.47	B_13
ATOM	2485	С	SER	108	69.645	68.260	-5.745	1.00 17.68	B_13
MOTA	2486	ŏ	SER	108	68.964	67.618	-6.541	1.00 19.67	D_13
									B_13
MOTA	2487	N	LYS	109	70.895	67.919	-5.448	1.00 11.70	B_13
MOTA	2489	CA	LYS	109	71.468	66.721	-6.047	1.00 10.00	B_13
MOTA	2490	CB	LYS	109	72.994	66.748	-5.989	1.00 18.86	B_13
ATOM	2491	CG	LYS	109	73.657	65.833	-7.013		
								1.00 16.33	B_13
MOTA	2492	CD	LYS	109	75.143	65.726	-6.740	1.00 11.58	B_13
ATOM	2493	CE	LYS	109	75.787	64.655	-7.606	1.00 27.43	B_13
ATOM	2494	NZ	LYS	109	77.218	64.492	-7.251	1.00 35.03	B_13
MOTA	2498	C	LYS	109	70.916	65.428	-5.444	1.00 29.39	B_13
MOTA	2499	0	LYS	109	71.432	64.905	-4.449	1.00 29.95	B_13
ATOM	2500	N	GLY	110	69.852	64.922	-6.055	1.00 14.77	B_13
ATOM	2502	CA	GLY	110	69.227	63.705	-5.576	1.00 24.08	B_13
MOTA	2503	С	GLY	110	67.793	64.105	-5.342	1.00 20.25	B_13
		ŏ							
ATOM	2504	-	GLY	110	67.203	64.737	-6.198	1.00 16.21	B_13
ATOM	2505	N	TYR	111	67.248	63.772	-4.182	1.00 10.00	B_13
MOTA	2507	CA	TYR	111	65.879	64.130	-3.845	1.00 24.52	B_13
MOTA	2508	CB	TYR	111	65.030	62.868	-3.688	1.00 22.46	B_13
ATOM	2509								
		CG	TYR	111	64.676	62.244	-4.999	1.00 10.83	B_13
MOTA	2510		TYR	111	65,380	61.155	-5.483	1.00 25.38	B_13
MOTA	2511	CE1	TYR	111	65.068	60.592	-6.720	1.00 18.68	B_13
ATOM	2512	CD2	TYR	111	63.646	62.769	-5.776	1.00 16.02	B 13
MOTA	2513	CE2	TYR	111	63.328	62.223	-7.013		
								1.00 31.72	B_13
MOTA	2514	CZ	TYR	111	64.041	61.131	-7.473	1.00 23.68	B_13
MOTA	2515	ОН	TYR	111	63.711	60.550	-8.666	1.00 20.96	B_13
ATOM	2517	С	TYR	111	65.856	64.944	-2.553	1.00 22.83	B_13
MOTA	2518	0	TYR	111	66.410	64.518	-1.538	1.00 11.66	B_13
ATOM	2519	N		112					
			ASN		65.278	66.140	-2.611	1.00 17.47	B_13
MOTA	2521	CA	ASN	112	65.180	67.006	-1.431	1.00 15.77	B_13
MOTA	2522	CB	ASN	112	64.658	68.401	-1.817	1.00 15.93	B_13
MOTA	2523	CG	ASN	112	64.694	69.384	-0.657	1.00 10.00	B_13
MOTA	2524		ASN	112	63.757	69.465	0.132	1.00 15.33	B_13
ATOM	2525								
	2323		ASN	112	65.754	70.180	-0.586	1.00 13.70	B_13
MOTA	2528	C	ASN	112	64.214	66.329	-0.472	1.00 17.73	B_13
ATOM	2529	0	ASN	112	63.007	66.243	-0.737	1.00 12.61	B_13
ATOM	2530	N	LEU	113	64.755	65.830	0.630	1.00 16.28	B_13
ATOM	2532	CA	LEU	113	63.962				
						65.121	1.619	1.00 15.93	B_13
MOTA	2533	CB	LEU	113	64.841	64.703	2.804	1.00 11.93	B_13
MOTA	2534	CG	LEU	113	64.719	63.352	3.521	1.00 17.15	B_13
MOTA	2535	CD1	LEU	113	65.002	63.640	4.987	1.00 10.00	B_13
ATOM	2536		LEU	113	63.370	62.667	3.362		5-13
								1.00 16.08	B_13
MOTA	2537	C	LEU	113	62.802	65.994	2.085	1.00 14.61	B_13
MOTA	2538	0	LEU	113	61.673	65:528	2.161	1.00 17.98	B_13
MOTA	2539	N	PHE	114	63.073	67.267	2.346	1.00 16.81	B_13
MOTA	2541	CA	PHE	114	62.056	68.212	2.791		
								1.00 15.65	B_13
MOTA	2542	CB	PHE	114	62.638	69.630	2.888	1.00 22.16	B_13
MOTA	2543	CG	PHE	114	61.596	70.714	2.882	1.00 12.27	B_13
MOTA	2544	CD1	PHE	114	60.804	70.952	4.004	1.00 19.93	B_13
MOTA	2545		PHE	114	61.378	71.470	1.746	1.00 13.56	B_13
ATOM	2546		PHE	114					
					59.813	71.932	3.984	1.00 17.08	B_13
ATOM	2547	CE2		114	60.398	72.441	1.726	1.00 13.79	B_13
MOTA	2548	CZ	PHE	114	59.615	72.666	2.848	1.00 10.70	B_13
MOTA	2549	С	PHE	114	60.860	68.220	1.842	1.00 19.55	B_13
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ATOM	2550	0	PHE	114	59.714	68.156	2.285	1.00 15.97	B_13
ATCM	2551		LEU	115	61.135	68.309	0.543	1.00 13.35	B_13
MOTA	2553		LEU	115	60.096	68.323	-0.485	1.00 17.91	B_13
MOTA	2554	CB	LEU	115	60.741	68.462	-1.868	1.00 24.65	B_13
ATOM	2555	CG	LEU	115	60.501	69.739	-2.679	1.00 22.70	B_13
ATOM	2556	CD1		115	61.033	70.939	-1.943	1.00 17.98	B_13
MOTA	2557	CD2	LEU	115	61.148	69.624	-4.048	1.00 28.50	B_13
MOTA	2558	С	LEU	115	59.235	67.042	-0.443	1.00 21.61	B_13
MOTA	2559	0	LEU	115	58.002	67.093	-0.344	1.00 13.99	B_13
MOTA	2560	N	VAL	116	59.898	65.895	-0.511	1.00 11.14	B_13
MOTA	2562	CA	VAL	116	59.199	64.616	-0.482	1.00 22.27	B_13
MOTA	2563	CB	VAL	116 ·	60.163	63.421	-0.772	1.00 17.40	B_13
ATOM	2564	CG1		116	59.437	62.086	-0.629	1.00 23.09	B_13
ATOM	2565	CG2	VAL	116	60.741	63.534	-2.169	1.00 12.16	B_13
ATOM	2566	С	VAL	116	58.502	64.414	0.864	1.00 10.00	B_13
ATOM	2567	Ō	VAL	116	57.368	63.950	0.911	1.00 16.18	B_13
ATOM	2568	N	ALA	117	59.153	64.803	1.954	1.00 10.00	B_13
ATOM	2570	CA	ALA	117	58.585	64.640	3.297	1.00 19.50	B_13
ATOM	2571	CB	ALA	117	59.608	64.995	4.352	1.00 11.81	B_13
ATOM	2572	C	ALA	117	57.309	65.455	3.505	1.00 30.87	B_13
ATOM	2573	Ō	ALA	117	56.327	64.955	4.053	1.00 10.00	B_13
ATOM	2574	N	ALA	118	57.322	66.714	3.087	1.00 24.62	B_13
ATOM	2576	CA	ALA	118	56.140	67.553	3.222	1.00 20.76	B_13
ATOM	2577	CB	ALA	118	56.407	68.917	2.654	1.00 16.19	B_13
ATOM	2578	C	ALA	118	54.968	66.894	2.485	1.00 20.54	B_13
ATOM	2579	ō	ALA	118	53.843	66.889	2.981	1.00 22.12	B_13
ATOM	2580	N	HIS	119	55.255	66.315	1.321	1.00 10.00	B_13
ATOM	2582	CA	HIS	119	54.259	65.647	0.489	1.00 17.27	B_13
ATOM	2583	CB	HIS	119	54.909	65.263	-0.860	1.00 11.16	B_13
ATOM	2584	CG	HIS	119	54.006	64.530	-1.813	1.00 26.59	, B_13
MOTA	2585		HIS	119	53.377	63.335	-1.706	1.00 16.63	B_13
ATOM	2586		HIS	119	53.723	64.995	-3.085	1.00 12.44	B_13
MOTA	2588		HIS	119	52.961	64.124	-3.715	1.00 14.58	B_13
ATOM	2589		HIS	119	52.734	63.101	-2.901	1.00 26.44	B_13
ATOM	2590	C	HIS	119	53.722	64.419	1.227	1.00 17.00	B_13
ATOM	2591	ŏ	HIS	119	52.510	64.218	1.331	1.00 17.01	B_13
MOTA	2592	N	GLU	120	54.626	63.607	1.751	1.00 10.31	B_13
MOTA	2594	CA	GLU	120	54.231	62.401	2.466	1.00 12.32	B_13
MOTA	2595	CB	GLU	120	55.463	61.627	2.961	1.00 15.34	B_13.
MOTA	2596	CG	GLU	120	56.354	61.078	1.848	1.00 10.00	B_13
ATOM	2597	CD	GLU	120	55.574	60.260	0.867	1.00 18.64	B_13
ATOM	2598		GLU	120	55.598	60.565	-0.348	1.00 18.04	B_13
ATOM	2599	OE2		120	54.920	59.308	1.320	1.00 18.08	B_13
ATOM	2600	C	GLU	120	53.347	62.777	3.635	1.00 12.41	B_13
ATOM	2601	o.	GLU	120	52.323	62.130	3.888	1.00 12.41	
ATOM	2602	N	PHE	121	53.750	63.813	4.359	1.00 20.02	B_13 B_13
MOTA	2604	CA	PHE	121	52.993	64.286	5.506		
MOTA	2605	CB	PHE	121	53.780	65.344	6.270	1.00 14.37 1.00 20.10	B_13 B_13
MOTA	2606	CG	PHE	121	55.057	64.827	6.852		B_13
ATOM	2607		PHE	121	56.037	65.700	7.292	1.00 24.55	B_13
ATOM	2608		PHE	121	55.292	63.454	6.936	1.00 23.62	B_13
ATOM	2609		PHE	121	57.247	65.212	7.813		B_13
MOTA	2610	CE2		121	56.488		7.448	1.00 18.59 1.00 15.21	
MOTA	2611	CZ			57.472	62.954	7.888		B_13 B_13
ATOM	2612	C	PHE	121 121	51.607	63.834 64.791	5.110	1.00 25.40	D-13
MOTA	2613	ŏ	PHE	121	50.676	64.760	5.921	1.00 16.63	B_13
ATOM	2614	N	GLY	122	51.471			1.00 26.80	B_13
ATOM	2616	CA	GLY	122	50.175	65.238 65.664	3.864 3.380	1.00 11.98 1.00 12.95	B_13
ATOM	2617	C	GLY	122	49.284			1.00 12.95	B_13
ATOM	2618			122		64.427	3.381		B_13
ATOM		0	GLY		48.113	64.483	3.753	1.00 13.74	B_13
ATOM	2619 2621	N	HIS		49.859	63.284	3.016	1.00 16.90	B_13
		CA	HIS	123	49.126	62.009	3.008	1.00 24.90	B_13
ATOM	2622	CB	HIS	123	49.918	60.918	2.279	1.00 18.28	B_13
ATOM	2623	CG	HIS	123	49.945	61.084	0.794	1.00 21.62	B_13
ATOM	2624		HIS	123	50.889	60.764	-0.119	1.00 13.04	B_13
ATOM	2625		HIS		48.887	61.618	0.093	1.00 17.18	B_13
ATOM	2627		HIS		49.176	61.621	-1.195	1.00 16.02	B_13
ATOM	2628		HIS	123	50.386	61.108	-1.353	1.00 15.58	B_13
MOTA	2629	C	HIS		48.864	61.562	4.446	1.00 19.74	B_13
ATOM	2630	0	HIS		47.744	61.179	4.785	1.00 15.41	B_13
MOTA	2631	N	SER		49.904	61.627	5.284	1.00 13.32	B_13
ATOM	2633	CA	SER		49.813	61.270	6.695		B_13
ATOM	2634	CB	SER		51.131	61.582	7.425	1.00 18.63	B_13
MOTA	2635	OG	SER		52.221	60.837	6.925	1.00 13.32	B_13
MOTA	2637	C	SER		48.703	62.102	7.335		B_13
MOTA	2638	0	SER		48.061	61.677	8.306		B_13
MOTA	2639	N	LEU	125	48.481	63.300	6.814	1.00 13.33	B_13

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ATOM	2641	CA	LEU	125	47.439	64.133	7.387	1.00 24.62	B_13
ATOM	2642	CB	LEU	125	47.893	65.592	7.436	1.00 20.76	B_13
MOTA	2643	CG	LEU	125	49.076	65.849	8.383	1.00 14.66	B_13
ATOM	2644	CD1		125	49.739		8.064		
						67.159		1.00 16.16	B_13
ATOM	2645	CD2		125	48.610	65.811	9.822	1.00 16.44	B_13
ATOM	2646	С	LEU	125	46.058	63.966	6.724	1.00 24.77	B_13
ATOM	2647	0	LEU	125	45.066	64.528	7.195	1.00 15.63	B_13
ATOM	2648	N	GLY	126					
					45.988	63.192	5.644	1.00 17.38	B_13
ATOM	2650	CA	GLY	126	44.700	62.968	5.001	1.00 22.41	B_13
ATOM	2651	C	GLY	126	44.453	63.487	3.603	1.00 13.20	B_13
ATOM	2652	٥	GLY	126					
					43.349	63.366	3.096	1.00 20.86	B_13
ATOM	2653	N	LEU	127	45.452	64.079	2.972	1.00 12.39	B_13
MOTA	2655	CA	LEU	127	45.267	64.592	1.617	1.00 11.56	B_13
ATOM	2656	СВ	LEU	127	45.965	65.947	1.467		
								1.00 19.19	B_13
MOTA	2657	CG	LEU	127	45.300	67.206	2.039	1.00 14.42	B_13
ATOM	2658	CD1	LEU	127	44.875	67.030	3.496	1.00 32.31	B_13
ATOM	2659	CD2	LEU	127	46.288	68.374	1.912	1.00 25.45	B_13
ATOM	2660	c	LEU	127					
					45.770	63.619	0.550	1.00 26.54	B_13
ATOM	2661	0	LEU	127	46.920	63.156	0.601	1.00 18.76	B_13
ATOM	2662	N	ASP	128	44.908	63.285	-0.407	1.00 28.54	B_13
ATOM	2664	CA	ASP	128	45.292	62.376	-1.480		
								1.00 10.89	B_13
ATOM	2665	CB	ASP	128	44.059	61.762	-2.136	1.00 15.95	B_13
ATOM	2666	CG	ASP	128	44.351	60.430	-2.794	1.00 23.44	B_13
MOTA	2667	OD1	ASP	128	43.377	59.735	-3.164	1.00 41.43	B_13
ATOM	2668		ASP	128					
					45.541	60.059	-2.918	1.00 18.12	B_13
ATOM	2669	С	ASP	128	46.060	63.203	-2.502	1.00 25.34	B_13
MOTA	2670	0	ASP	128	46.489	64.308	-2.213	1.00 16.36	B_13
MOTA	2671	N	HIS	129	46.283		-3.682		
						62.645		1.00 17.53	B_13
MOTA	2673	CA	HIS	129	47.001	63.366	-4.718	1.00 26.87	B_13
ATOM	2674	CB	HIS	129	47.495	62.398	-5.794	1.00 10.00	B_13
MOTA	2675	CG	HIS	129	48.729	61.645			
							-5.400	1.00 19.64	B_13
MOTA	2676		HIS	129	49.769	61.996	-4.609	1.00 19.96	B_13
MOTA	2677	ND1	HIS	129	49.012	60.373	-5.859	1.00 23.97	B_13
ATOM	2679	CEI	HIS	129	50.170	59.977	-5.372	1.00 17.95	B_13
ATOM	2680		HIS	129					
					50.658	60.944	-4.605	1.00 13.79	B_13
MOTA	2681	С	HIS	129	46.153	64.457	-5.360	1.00 39.97	B_13
MOTA	2682	0	HIS	129	45.011	64.220	-5.757	1.00 25.97	B_13
ATOM	2683	N	SER	130	46.743	65.640	-5.481		
								1.00 21.04	B_13
ATOM	2685	CA	SER	130	46.090	66.776	-6.109	1.00 16.72	B_13
ATOM	2686	ÇВ	SER	130	46.847	68.058	-5.757	1.00 20.97	B_13
ATOM	2687	OG	SER	130	46.358	69.154	-6.502	1.00 25.52	
	2689								B_13
ATOM		C	SER	130	46.098	66.582	-7.622	1.00 24.66	B_13
MOTA	2690	0	SER	130	46.779	65.694	-8.145	1.00 29.24	B_13
ATOM	2691	N	LYS	131	45.315	67.403	-8.315	1.00 26.96	B_13
ATOM	2693	CA	LYS	131					
					45.253	67.358	-9.769	1.00 20.25	B_13
MOTA	2694	CB	LYS	131	43.796	67.379	-10.247	1.00 33.22	B_13
MOTA	2695	CG	LYS	131	43.159	68.775	-10.302	1.00 32.85	B_13
ATOM	2696	CD	LYS	131	43.335				
ATOM							-11.675	1.00 15.99	B_13
	2697	CE	LYS	131	43.023		-11.601	1.00 30.34	B_13
ATOM	2698	NZ	LYS	131	43.879	71.647	-10.600	1.00 30.44	B_13
ATOM	2702	С	LYS	131	45.998	68.602	-10.249	1.00 15.31	B_13
ATOM	2703	ō	LYS	131					
					46.414		-11.402	1.00 30.72	B_13
ATOM	2704	N	ASP	132	46.191	69.536	-9.323	1.00 23.41	B_13
MOTA	2706	CA	ASP	132	46.869	70.798	-9.581	1.00 22.69	B 13
MOTA	2707	CB	ASP	132	46.641	71.726	-8.379	1.00 24.86	B_13
ATOM	2708	CG	ASP	132	46.819	73.200			
							-8.712	1.00 24.93	B_13
ATOM	2709		ASP	132	46.007	74.009	-8.208	1.00 29.71	B_13
MOTA	2710	OD2	ASP	132	47.766	73.555	-9.448	1.00 28.82	B_13
ATOM	2711	С	ASP	132	48.358	70.497	-9.728	1.00 14.97	B_13
ATOM	2712	ō	ASP	132					
					49.047	70.235	-8.742	1.00 19.64	B_13
MOTA	2713	N	PRO	133	48.874	70.538	-10.964	1.00 16.94	B_13
MOTA	2714	CD	PRO	133	48.209		-12.199	1.00 21.42	B_13
MOTA	2715	CA	PRO	133					5_13
					50.293		-11.215	1.00 19.34	B_13
MOTA	2716	CB	PRO	133	50.457	70.636	-12.690	1.00 20.48	B_13
MOTA	2717	CG	PRO	133	49.347		-12.929	1.00 21.80	B_13
MOTA	2718	c	PRO	133	51.237				
							-10.322	1.00 17.45	B_13
ATOM	2719	0	PRO	133	52.319	70.590		1.00 23.30	B_13
MOTA	2720	N	GLY	134	50.799	72.246	-9.904	1.00 32.46	B_13
ATOM	2722	CA	GLY	134	51.610	73.104	-9.051	1 00 10 44	
ATOM	2723				21.010			1.00 19.44	B_13
		Ç	GLY	134	51.306	72.958	-7.569	1.00 22.33	B_13
ATOM	2724	0	GLY	134	51.556	73.877	-6.795	1.00 21.92	B_13
ATOM	2725	N	ALA	135	50.698	71.836	-7.190	1.00 34.71	B_13
ATOM	2727	CA	ALA	135					
					50.355	71.580	-5.794	1.00 18.35	B_13
MOTA	2728	CB	ALA	135	48.948	70.987	-5.690	1.00 14.30	B_13
MOTA	2729	С	ALA	135	51.370	70.616	-5.210	1.00 10.00	B_13
ATOM	2730	0	ALA	135	51.739				
ATOM	2731	N	LEU	136		69.647	-5.858	1.00 17.52	B_13
011	-,51	*4	LEU	430	51.727	70.842	-3.952	1.00 21.29	B_13

MOTA	2733		LEU	136	52.692	70.015 -3.230	1.00 14.62	B_13
MOTA	2734		LEU	136	52.738	70.458 -1.763	1.00 18.54	B_13
ATOM	2735	CG	LEU	136	54.007	70.308 -0.921	1.00 34.11	B_13
MOTA	2736	CD1	LEU	136	53.587	69.907 0.485	1.00 14.76	B_13
ATOM	2737	CD2	LEU	136	54.969	69.296 -1.508	1.00 11.64	B_13
ATOM	2738	С	LEU	136	52.232	68.564 -3.287	1.00 13.50	B_13
MOTA	2739		LEU	136	53.033	67.640 -3.238	1.00 19.04	B_13
ATOM	2740		MET	137	50.921	68.364 -3.281	1.00 17.54	B_13
ATOM	2742		MET	137	50.360	67.019 -3.324		
MOTA	2743		MET	137	49.010		1.00 25.11	B_13
	2744					66.981 -2.599	1.00 19.80	B_13
MOTA		CG	MET	137	49.083	67.312 -1.117	1.00 15.35	B_13
ATOM	2745	SD	MET	137	50.354	66.361 -0.262	1.00 11.22	B_13
ATOM	2746	CE	MET	137	49.882	64.680 -0.764	1.00 13.90	B_13
MOTA	2747	Ç	MET	137	50.254	66.387 -4.721	1.00 28.08	B_13
ATOM	2748	0	MET	137	49.730	65.268 -4.863	1.00 12.18	B_13
MOTA	2749	N	PHE	138	50.771	67.070 -5.743	1.00 10.00	B_13
ATOM	2751	CA	PHE	138	50.751	66.528 -7.097	1.00 12.27	B_13
MOTA	2752	CB	PHE	138	51.327	67.523 -8.094	1.00 19.38	B_13
MOTA	2753	CG	PHE	138	51.051	67.175 -9.534	1.00 25.74	B_13
MOTA	2754	CD1	PHE	138	52.090	67.077 -10.448	1.00 19.74	B_13
ATOM	2755	CD2	PHE	138	49.747	67.007 -9.990	1.00 24.46	B_13
ATOM	2756	CE1	PHE	138	51.843	66.824 -11.786	1.00 19.54	B_13
ATOM	2757	CE2		138	49.495	66.750 -11.335	1.00 24.12	B_13
ATOM	2758	CZ	PHE	138	50.544	66.664 -12.230	1.00 18.15	B_13
ATOM	2759	c	PHE	138	51.619	65.269 -7.068	1.00 25.93	B_13
ATOM	2760	ŏ	PHE	138	52.658	65.226 -6.414	1.00 23.93	
ATOM	2761	N	PRO	139	51.166	64.194 -7.714		B_13
MOTA	2762	CD			49.870		1.00 25.17	B_13
			PRO	139		64.004 -8.392	1.00 10.00	B_13
MOTA	2763	CA	PRO	139	51.950	62.956 -7.713	1.00 18.48	B_13
MOTA	2764	CB	PRO	139	50.981	61.946 -8.339	1.00 15.96	B_13
MOTA	2765	CG	PRO	139	50.140	62.798 -9.250	1.00 18.82	B_13
MOTA	2766	C	PRO	139	53.299	62.950 -8.430	1.00 17.22	B_13
ATOM	2767	0	PRO	139	53.849	61.876 -8.661	1.00 36.93	B_13
MOTA	2768	N	ILE	140	53.844	64.114 -8.767	1.00 24.48	B_13
MOTA	2770	CA	ILE	140	55.118	64.155 -9.477	1.00 20.03	B_13
MOTA	2771	CB	ILE	140	54.996	64.807 -10.892	1.00 18.71	B_13
MOTA	2772	CG2	ILE	140	56.334	64.709 -11.639	1.00 23.96	B_13
ATOM	2773	CG1	ILE	140	53.932	64.113 -11.724	1.00 24.68	B_13
ATOM	2774		ILE	140	53.861	64.669 -13.125	1.00 25.83	B_13
ATOM	2775	C	ILE	140	56.109	64.992 -8.700	1.00 27.87	B_13
ATOM	2776	ō	ILE	140	55.758	66.043 -8.248	1.00 22.39	B_13
ATOM	2777	N	TYR	141	57.332	64.512 -8.535	1.00 12.36	
ATOM	2779	CA	TYR	141	58.350	65.281 -7.834		B_13
MOTA	2780	CB	TYR	141	59.418		1.00 21.85	B_13
ATOM	2781	CG	TYR	141		64.353 -7.266	1.00 15.16	B_13
ATOM	2782		TYR		60.592	65.096 -6.672	1.00 15.65	B_13
				141	61.755	65.306 -7.407	1.00 18.56	B_13
MOTA	2783	CE1		141	62.836	65.967 -6.859	1.00 10.00	B_13
ATOM	2784	CD2	TYR	141	60.546	65.576 -5.366	1.00 11.42	B_13
ATOM	2785	CE2	TYR	141	61.626	66.236 -4.814	1.00 13.45	B_13
ATOM	2786	CZ	TYR	141	62.770	66.429 -5.567	1.00 10.00	B_13
MOTA	2787	OH	TYR	141	63.841	67.109 -5.016	1.00 18.97	B_13
MOTA	2789	С	TYR	141	59.042	66.270 -8.776	1.00 19.52	B_13
ATOM	2790	0	ŢYR	141	59.709	65.859 -9.727	1.00 21.37	B_13
ATOM	2791	N	THR	142	58.932	67.556 -8.465	1.00 23.99	B_13
ATOM	2793	CA	THR	142	59.573		1.00 19.53	B_13
MOTA	2794	CB	THR	142	58.515	69.578 -9.807	1.00 10.00	B_13
MOTA	2795		THR	142	57.704	68.880 -10.756	1.00 37.02	B_13
MOTA	2797		THR	142	59.151	70.757 -10.457	1.00 34.35	B_13
MOTA	2798	С	THR	142	60.483	69.332 -8.235	1.00 19.89	B_13
ATOM	279 <del>9</del>	0	THR	142	60.120	69.513 -7.076	1.00 25.67	B_13
ATOM	2800	N	TYR	143	61.699	69.677 -8.643	1.00 30.64	B_13
MOTA	2802	CA	TYR	143	62.609	70.344 -7.707	1.00 32.54	B_13
ATOM	2803	CB	TYR	143	64.091	70.190 -8.108	1.00 26.34	B_13
ATOM	2804	CG	TYR	143	65.008	71.048 -7.244	1.00 10.69	B_13
ATOM	2805		TYR	143	65.066	70.866 -5.852	1.00 16.37	B_13
ATOM	2806	CE1		143	65.801	71.738 -5.035	1.00 16.37	D 13
ATOM	2807	CD2		143				B_13
MOTA	2808	CE2			65.714	72.114 -7.795	1.00 17.36	B_13
				143	66.451	73.006 -6.981	1.00 15.32	B_13
MOTA	2809	CZ	TYR	143	66.489	72.810 -5.610	1.00 10.00	B_13
MOTA	2810	ОН	TYR	143	67.184	73.665 -4.790	1.00 27.84	B_13
MOTA	2812	C	TYR	143	62.330	71.815 -7.456	1.00 24.77	B_13
ATOM	2813	0	TYR	143	62.201	72.611 -8.399	1.00 26.19	B_13
ATOM	2814	N	THR	144	62.292	72.160 -6.170	1.00 22.23	B_13
ATOM	2816	CA	THR	144	62.103	73.533 -5.727	1.00 33.68	B_13
MOTA	2817	CB	THR	144	60.668	73.814 -5.189	1.00 28.06	B_13
ATOM	2818		THR	144	60.277	72.812 -4.241	1.00 38.14	B_13
MOTA	2820	CG2	THR	144	59.681	73.857 -6.346	1.00 48.73	B_13
								_

ATOM	2021	_	mun.	1 4 4	62 120	72 002	4 605		
	2821	С	THR	144	63.178	73.893	-4.695	1.00 35.52	B_13
MOTA	2822	0	THR	144	64.207	74.465	-5.064	1.00 39.57	B_13
ATOM	2823	N	GLY	145	62.967	73.552	-3.422	1.00 35.95	
		-							B_13
MOTA	2825	CA	GLY	145	63.967	73.872	-2.407	1.00 35.01	B_13
ATOM	2826	С	GLY	145	63.509	74.025	-0.965	1.00 26.81	B_13
ATOM									
	2827	0	GLY	145	62.566	74.773	-0.670	1.00 40.81	B_13
MOTA	2828	N	LYS	146	64.302	73.439	-0.066	1.00 27.13	B_13
MOTA	2830	CA	LYS	146	64.071	73.423	1.389		
								1.00 23.89	B_13
MOTA	2831	CB	LYS	146	65.163	72.548	2.049	1.00 29.08	B_13
ATOM	2832	CG	LYS	146	64.992	72.209	3.524	1.00 19.99	B_13
ATOM	2833	CD	LYS	146	66.079	71.224	3.913	1.00 20.44	B_13
ATOM	2834	CE	LYS	146	66.181	71.010	5.402	1.00 24.16	B_13
ATOM	2835	NZ	LYS	146	67.250				5-13
						69.987	5.727	1.00 23.37	B_13
MOTA	2839	С	LYS	146	63.926	74.778	2.124	1.00 18.98	B_13
ATOM	2840	0	LYS	146	63.900	74.831	3.353	1.00 28.15	
									B_13
MOTA	2841	N	SER	147	63.826	75.871	1.382	1.00 35.50	B_13
MOTA	2843	CA	SER	147	63.661	77.185	1.992	1.00 31.59	B_13
ATOM	2844								
		CB	SER	147	64.988	77.673	2.594	1.00 27.05	B_13
MOTA	2845	OG	SER	147	65.996	77.756	1.586	1.00 48.28	B_13
ATOM	2847	С	SER	147	63.203	78.131	0.902		
								1.00 27.12	B_13
MOTA	2848	0	SER	147	62.743	79.251	1.168	1.00 33.75	B_13
MOTA	2849	N	HIS	148	63.248	77.644	-0.332	1.00 25.13	B_13
ATOM	2851	CA	HIS						
				148	62.872	78.465	-1.463	1.00 23.42	B_13
MOTA	2852	CB	HIS	148	63.704	78.076	-2.678	1.00 17.40	B_13
MOTA	2853	ÇG	HIS	148	65.174	78.020	-2.398	1.00 45.97	B_13
MOTA	2854	CD2	HIS	148	66.204	77.524	-3.121	1.00 27.24	B_13
MOTA	2855	ND1	HIS	148	65.724	78.476	-1.213	1.00 43.49	B_13
ATOM	2857		HIS						
				148	67.024	78.253	-1.218	1.00 30.28	B_13
MOTA	2858	NE2	HIS	148	67.342	77.676	-2.366	1.00 45.28	B_13
ATOM	2860	С	HIS	148	61.381	78.433	-1.796	1.00 47.15	B_13
							-1.750		
MOTA	2861	0	HIS	148	60.936	79.166	-2.704	1.00 40.97	B_13
ATOM	2862	N	PHE	149	60.601	77.636	-1.053	1.00 48.76	B_13
ATOM	2864	CA	PHE	149					
					59.170	77.557	-1.347	1.00 32.44	B_13
MOTA	2865	CB	DHE.	149	58.856	76.364	-2.269	1.00 27.77	B_13
ATOM	2866	CG	PHE	149	58.415	76.781	-3.657	1.00 24.63	B_13
MOTA	2867		PHE	149	57.826	75.874	-4.520	1.00 25.66	B_13
MOTA	2868	CD2	PHE	149	58.550	78.106	-4.072	1.00 30.89	B_13
MOTA	2869		PHE						
				149	57.376	76.277	-5.767	1.00 17.10	B_13
ATOM	2870	CE2	PHE	149	58.104	78.520	-5.311	1.00 18.57	B_13
MOTA	2871	CZ	PHE	149	57.513	77.608	-6.166	1.00 30.20	B_13
									P_T2
MOTA	2872	С	PHE	149	58.061	77.791	-0.308	1.00 27.40	B_13
ATOM	2873	0	PHE	149	58.299	77.971	0.892	1.00 29.69	B_13
ATOM	2874	N	MET						
				150	56.836	77.729	-0.822	1.00 28.66	B_13
ATOM	2876	ÇA	MET	150	55.621	78.027	-0.094	1.00 20.63	B_13
MOTA	2877	CB	MET	150	55.251	79.431	-0.503	1.00 25.60	B_13
									B_13
MOTA	2878	ÇG	MET	150	55.599	79.691	-1.989	1.00 23.95	B_13
ATOM	2879	SD	MET	150	57.336	80.086	-2.296	1.00 76.68	B_13
MOTA	2880	CE	MET	150	57.209	81.473	-3.385		
								1.00 21.07	B_13
ATOM	2881	С	MET	150	54.436	77.118	-0.450	1.00 30.58	B_13
ATOM	2882	0	MET	150	54.104	76.948	-1.628	1.00 16.91	B_13
ATOM	2883	N	LEU	151					
					53.727	76.664	0.581	1.00 36.94	B_13
MOTA	2885	CA	LEU	151	52.576	75.772	0.431	1.00 25.68	B_13
MOTA	2886	CB	LEU	151	51.968	75.474	1.807		B 13
ATOM	2887								
		CG	LEU	151	51.087	74.232	1.927	1.00 24.21	B_13
MOTA	2888		LEU	151	51.936	72.998	1.657	1.00 21.54	_B_13
ATOM	2889		LEU	151	50.487	74.150	3.314	1.00 19.89	B_13
ATOM									5-13
	2890	C	LEU	151	51.498	76.322	-0.491	1.00 17.09	B_13
ATOM	2891	0	LEU	151	50.795	77.267	-0.136	1.00 35.38	B_13
MOTA	2892	N	PRO	152	51.338	75.727	-1.686		
								1.00 16.90	B_13
ATOM	2893	CD	PRO	152	52.154	74.643	-2.255	1.00 25.80	B_13
ATOM	2894	CA	PRO	152	50.334	76.170	-2.653	1.00 29.65	B_13
ATOM	2805	CB	PRÒ	152					2-13
	2895 2896				50.447	75.110	-3.749	1.00 24.68	B_13
MOTA	2896	CG	PRO	152	51.892	74.791	-3.722	1.00 14.34	B_13
MOTA	2897	С	PRO	152	48.910	76.261	-2.087	1.00 10.00	B_13
ATOM	2898	0	PRO	152	48.543	75.505	-1.184	1.00 20.25	B_13
MOTA	2899	N	ASP	153	48.117	77.180	-2.639	1.00 19.53	B_13
ATOM	2901	CA	ASP	153					
					46.723	77.387	-2.226	1.00 15.90	B_13
ATOM	2902	CB	ASP	153	45.986	78.304	-3.213	1.00 22.34	B_13
MOTA	2903	CG	ASP	153	46.418	79.741	-3.115		B_13
								1.00 28.86	
ATOM	2904		ASP	153	47.016	80.115	-2.074	1.00 35.34	B_13
MOTA	2905	OD2	ASP	153	46.142	80.494	-4.084	1.00 30.09	B_13
ATOM	2906		ASP	153					
		c			45.953	76.084	-2.169	1.00 27.31	B_13
MOTA	2907	0	ASP	153	45.309	75.783	-1.167	1.00 23.50	B_13
ATOM	2908	N	ASP	154	46.000	75.339	-3.276	1.00 25.51	B_13
ATOM									
	2910	CA	ASP	154	45.316	74.063	-3.392	1.00 20.91	B_13
ATOM	2911	CB	ASP	154	45.745	73.364	-4.682	1.00 14.23	B_13
ATOM	2912	CG	ASP	154	45.033	72.062	-4.885	1.00 22.95	B_13
	~						-4.000	1.00 66.33	2_13

ATOM	2913	OD1	ACD	154	45.590	71.026	-4.516	1.00 17.80	B_13
ATOM	2914	OD2		154	43.904	72.076	-5.388	1.00 19.14	B_13
MOTA	2915		ASP	154	45.551	73.155	-2.173	1.00 26.95	B_13
ATOM	2916		ASP	154	44.629	72.491	-1.696	1.00 22.92	B_13
MOTA MOTA	2917		ASP	155	46.776 47.110	73.155	-1.654 -0.490	1.00 23.56 1.00 28.69	B_13 B_13
ATOM	2919 2920		ASP ASP	155 155	48.618	72.338 72.118	-0.388	1.00 28.89	B_13 B_13
MOTA	2921		ASP	155	49.208	71.566	-1.676	1.00 24.35	B_13
ATOM	2922	OD1		155	49.705	72.369	-2.500	1.00 27.89	B_13
MOTA	2923	OD2		155	49.152	70.335	-1.875	1.00 16.96	B_13
MOTA	2924	c	ASP	155	46.582	72.976	0.781	1.00 25.41	B_13
MOTA	2925 2926	0 N	ASP	155	46.055 46.733	72.275 74.296	1.656 0.891	1.00 13.36 1.00 16.99	B_13
ATOM ATOM	2928		VAL VAL	156 156	46.222	75.021	2.053	1.00 22.26	B_13 B_13
ATOM	2929	СВ	VAL	156	46.340	76.571	1.901	1.00 25.69	B_13
MOTA	2930	CG1		156	45.811	77.249	3.158	1.00 14.95	B_13
MOTA	2931	CG2		156	47.768	77.007	1.641	1.00 17.52	B_13
MOTA	2932	C	VAL	156	44.727	74.705	2.129	1.00 10.00	B_13
MOTA MOTA	2933 2934	N O	VAL GLN	156 157	44.224 44.033	74.234 74.980	3.145 1.029	1.00 22.47 1.00 16.19	B_13 B_13
ATOM	2936	CA	GLN	157	42.604	74.758	0.930	1.00 17.97	B_13
MOTA	2937	CB	GLN	157	42.108	75.039	-0.497	1.00 17.10	B_13
MOTA	2938	CG	GLN	157	40.804	75.852	-0.547	1.00 26.00	B_13
MOTA	2939	CD	GLN	157	40.949	77.284	-0.005	1.00 25.84	B_13
MOTA	2940 2941	OE1 NE2	GLN GLN	157 157	41.218 40.744	77.505 78.255	1.177 -0.875	1.00 39.61 1.00 32.22	B_13 B_13
MOTA MOTA	2944	C	GLN	157	42.347	73.324	1.309	1.00 18.69	B_13
MOTA	2945	ŏ	GLN	157	41.368	73.015	1.982	1.00 10.00	B_13
MOTA	2946	N	GLY	158	43.272	72.460	0.903	1.00 31.05	B_13
MOTA	2948	CA	GLY	158	43.156	71.053	1.205	1.00 21.69	B_13
MOTA	2949	C	GLY	158	43.129	70.738	2.684	1.00 13.51	B_13
MOTA MOTA	2950 2951	о И	GLY ILE	158 159	42.108 44.224	70.263 71.006	3.182 3.398	1.00 14.91 1.00 19.34	B_13 B_13
MOTA	2953	CA	ILE	159	44.268	70.686	4.827	1.00 19.14	B_13
MOTA	2954	CB	ILE	159	45.669	70.880	5.503	1.00 12.57	B_13
MOTA	2955		ILE	159	46.268	69.542	5.960	1.00 19.22	B_13
MOTA	2956		ILE	159	46.603	71.702	4.633	1.00 31.62	B_13
MOTA	2957		ILE	159	46.426 43.235	73.177 71.461	4.824 5.610	1.00 25.87 1.00 21.87	B_13 B_13
MOTA MOTA	2958 2959	С.	ILE	159 159	42.691	70.952	6.592	1.00 21.07	B_13
MOTA	2960	Ŋ	GLN	160	42.959	72.689	5.186	1.00 12.08	B_13
ATOM	2962	CA	GLN	160	41.967	73.483	5.874	1.00 11.43	B_13
MOTA	2963	CB	GLN	160	41.949	74.916	5.346	1.00 29.25	B_13
ATOM	2964	CG	GLN	160	43.158	75.737	5.827	1.00 22.01	B_13
ATOM ATOM	2965 2966	CD OE1	GLN GLN	160 160	43.098 42.260	77.199 77.593	5.416 4.607	1.00 18.77 1.00 36.02	B_13 B_13
MOTA	2967	NE2		160	43.997	78.004	5.965	1.00 28.49	B_13
ATOM	2970	C	GLN	160	40.596	72.820	5.772	1.00 22.28	B_13
ATOM	2971	0	GLN	160	39.855	72.786	6.754	1.00 14.16	B_13
MOTA	2972	N	SER	161	40.304	72.183	4.634	1.00 32.89	B_13
ATOM ATOM	2974 2975	CA CB	SER SER	161 161	39.005 38.847	71.537 70.901	4.474	1.00 29.25 1.00 19.70	B_13 B_13
ATOM	2976	OG	SER	161	39.594	69.706	2.946	1.00 24.88	B_13
MOTA	2978	C	SER	161	38.831	70.503	5.566	1.00 22.08	B_13
MOTA	2979	0	SER	161	37.745	70.340	6.118	1.00 26.26	B_13
MOTA MOTA	2980 2982	N CA	LEU	162 162	39.931 39.913	69.852 68.829	5.919 6.953	1.00 19.14 1.00 29.17	B_13 B_13
MOTA	2983	CB	LEU	162	41.081	67.852	6.767	1.00 12.08	B_13
MOTA	2984	CG	LEU	162	40.982	66.666	5.812	1.00 20.09	B_13
ATOM	2985		. LEU	162	40.661	67.184	4.478	1.00 24.51	B_13
MOTA	2986		LEU	162	42.299	65.884	5.794	1.00 27.00	B_13
MOTA MOTA	2987 2988	C O	LEU LEU	162 162	39.965 39.047	69.392 69.191	8.364 9.162	1.00 24.75 1.00 22.04	B_13 B_13
ATOM	2989	N	TYR	163	41.015	70.151	8.652	1.00 20.72	B_13
ATOM	2991	CA	TYR	163	41.211	70.689	9.980	1.00 10.00	B_13
MOTA	2992	CB	TYR	163	42.695	70.595	10.343	1.00 10.95	B_13
MOTA	2993	CG	TYR	163	43.221	69.167	10.209		B_13
MOTA	2994	CD1		163 163	43.114	68.261	11.264	1.00 37.53	B_13
MOTA MOTA	2995 2996	CE1	L TYR	163 163	43.452 43.703	66.913 68.689	11.103 8.990	1.00 26.00 1.00 23.78	B_13 B_13
ATOM	2997	CE2		163	44.048	67.342	8.822		B_13
MOTA	2998	CZ	TYR	163	43.914	66.461	9.879		B_13
MOTA	2999	OH	TYR	163	44.210	65.121	9.711	1.00 13.27	B_13
MOTA	3001	C	TYR	163	40.634	72.085			B_13
MOTA MOTA	3002 3003	N O.	TYR GLY	163 164	39.975 40.819	72.327 72.975			B_13 B_13
MOTA	3005		GLY		40.291				B_13
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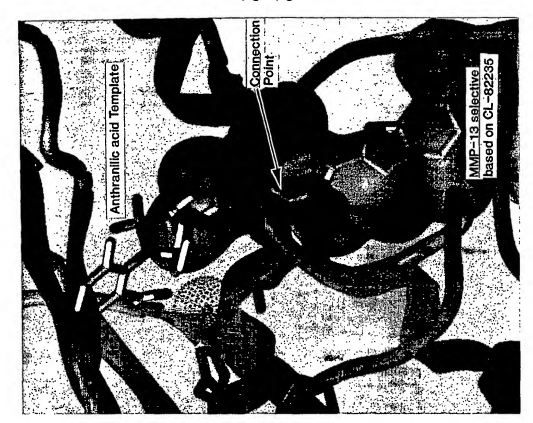
MOTA	3006	C GLY	164	41.402	75.344	9.424	1.00 30.89	B_13
ATOM	3C07	O GLY	164	41.101	76.564	9.368	1.00 26.89	B_13
MOTA	3008	OT GLY	164	42.570	74.911	9.560	1.00 27.71	
								B_13
MOTA	3013	ZN ZN	166	51.961	60.891	-2.865	1.00 28.31	BION
ATOM	3014	ZN ZN	167	56.468	50.981	3.458	1.00 26.20	BION
MOTA	3015	CA CA	168	63.096	53.752	-5.445	1.00 14.89	BION
MOTA	3016	CA CA	165	50.705	55.618	13.085	1.00 15.79	
								BION
MOTA	3047	C5 WAY	169	54.585	56.119	-6.288	1.00 40.09	B693
MOTA	3048	CF1 WAY	169	54.019	54.934	-5.802	1.00 21.52	B693
ATOM	3049	CH WAY	169	53.271	54.923	-4.624	1.00 32.32	
								B693
ATOM	3050	C2 WAY	169	53.100	56.104	-3.898	1.00 21.39	B693
ATOM	3051	C3 WAY	169	53.667	57.286	-4.369	1.00 18.26	B693
MOTA	3052	C4 WAY	169	54.402	57.308	-5.540	1.00 20.63	B693
MOTA	3053	N20 WAY	169	54.933	58.531	-5.964	1.00 22.15	B693
MOTA	3054	CD WAY	169	54.297	59.340	-7.031	1.00 30.92	B693
MOTA	3055	C23 WAY	169	53.576	58.491	-8.087	1.00 20.75	B693
ATOM	3056	C28 WAY	169					
				54.224	58.114	-9.279	1.00 34.14	B693
MOTA	3057	C27 WAY	169	53.539	57.335	-10.228	1.00 33.99	B693
ATOM	3058	CM WAY	169	52.209	56.944	-9.968	1.00 23.49	B693
ATOM	3059	N25 WAY	169	51.602	57.318	-8.814	1.00 23.61	B693
ATOM	3060	C24 WAY	169	52.246	58.071	-7.880	1.00 20.52	B693
MOTA	3061	S21 WAY	169	56.531	58.783	-5.660	1.00 20.46	B693
MOTA	3062	C16 WAY	169	56.457	60.446	-5.010	1.00 39.00	B693
ATOM	3063	C21 WAY	169	56.700	60.669		1.00 28.79	
						-3.634		B693
MOTA	3064	C20 WAY	169	56.656	61.967	-3.109	1.00 12.65	B693
MOTA	3065	C19 WAY	169	56.373	63.058	-3.946	1.00 15.68	B693
MOTA	3066	C18 WAY	169	56.126	62.828	-5.319	1.00 12.08	
								B693
ATOM	3067	C17 WAY		56.169	61.538	-5.852	1.00 15.19	B693
MOTA	3068	033 WAY	169	56.337	64.360	-3.424	1.00 16.79	B693
MOTA	3069	C36 WAY	169	56.982	65.456	-4.084	1.00 20.80	B693
ATOM	3070	O15 WAY		56.973	57.923	-4.580	1.00 21.90	B693
MOTA	3071	O14 WAY	169	57.259	58.799	-6.913	1.00 10.86	B693
ATOM	3072	C7 WAY	169	53.486	58.556	-3.613	1.00 10.00	B693
ATOM	3073	N9 WAY		53.741				
					58.606	-2.303	1.00 10.00	B693
ATOM	3074	010 WAY	169	53.539	59.846	-1.659	1.00 23.73	B693
ATOM	3075	OS WAY	169	53.107	59.569	-4.154	1.00 15.89	B693
ATOM	3076	C29 WAY		55.383	55.968	-7.606		
							1.00 28.30	B693
ATOM	1	OH2 WAT		67.399	53.332	19.612	1.00 10.00	SOLV
ATOM	2	OH2 WAT	302	61.288	46.506	17.898	1.00 10.00	SOLV
ATOM	3	OH2 WAT		79.538	50.433	20.115	1.00 10.00	
								SOLV
ATOM	4	OH2 WAT		80.982	25.236	19.076	1.00 26.37	SOLV
ATOM	5	OH2 WAT	305	82.461	30.767	19.346	1.00 13.02	SOLV
MOTA	6	OH2 WAT	306	67.759	41.912	4.887	1.00 17.30	SOLV
ATOM	7							
		OH2 WAT		60.785	41.727	10.585	1.00 20.42	SOLV
MOTA	8	OH2 WAT	, 308	89.638	33.523	25.640	1.00 33.45	SOLV
ATOM	9	OH2 WAT	309	77.721	51.975	4.391	1.00 13.91	SOLV
ATOM	10			96.022	34.702	6.692		
							1.00 25.50	SOLV
ATOM	11	OH2 WAT		71.292	38.746	26.741	1.00 13.06	SOLV
MOTA	12	OH2 WAI	312	85.939	49.781	3.498	1.00 12.04	SOLV
MOTA	13	OH2 WAT	313	58.101	41.127	10.261	1.00 40.97	SOLV
ATOM	14	OH2 WAT		86.373				
					42.692	0.747	1.00 17.24	SOLV
MOTA	15			78.257	39.885	24.626	1.00 18.57	SOLV
MOTA	16	OH2 WAT	316	68.341	48.572	25.558	1.00 18.33	SOLV
MOTA	17	OH2 WAT	317	79.806	29.147	18.371	1.00 10.00	SOLV
ATOM	18			87.119				
					44.480	23.137	1.00 46.31	SOLV
MOTA	19			55.885	39.688	11.459	1.00 21.26	SOLV
ATOM	20	OH2 WAT	320	73.250	41.084	0.386	1.00 18.49	SOLV
ATOM	21			72.079	46.488	-6.835	1.00 27.48	SOLV
MOTA	22			71.923	37.638	-3.750	1.00 29.19	SOLV
MOTA	23	OH2 WAT	323	74.998	28.451	2.684	1.00 34.60	SOLV
MOTA	24			87.769	44.123	9.214	1.00 15.60	SOLV
MOTA	25			86.113	24.382	16.709	1.00 25.17	SOLV
MOTA	26	OH2 WAT	326	81.205	57.603	0.529	1.00 34.27	SOLV
MOTA	27	OH2 WAT		75.163	62.739	12.391	1.00 16.47	SOLV
ATOM	28							
				65.604	44.690	2.830	1.00 26.64	SOLV
MOTA	29			61.899	45.512	29.269	1.00 15.82	SOLV
MOTA	30	OH2 WAT	330	58.763	41.730	8.338	1.00 27.95	SOLV
MOTA	31			69.823				
					44.729	6.258	1.00 13.37	SOLV
ATOM	32			79.220	61.263	12.781	1.00 28.84	SOLV
MOTA	33	OH2 WAT	333	78.105	37.095	27.911	1.00 34.48	SOLV
ATOM	34			75.939	25.608	12.364	1.00 35.21	SOLV
ATOM								
	35			90.256	42.668	16.539	1.00 45.05	SOLV
ATOM	36	OH2 WAT	336	86.761	51.457	13.881	1.00 25.26	SOLV
MOTA	37	OH2 WAT	337	67.479	42.004	-5.009	1.00 33.30	SOLV
ATOM	38							
				82.018	50.963	8.823	1.00 19.80	SOLV
ATOM	39			80.278	32.895	-1.126	1.00 30.16	SOLV
MOTA	40	OH2 WAT	r 340	71.683	50.944	31.567	1.00 29.62	SOLV
-							25.02	2024

ATOM	41	OH2 WAT	341	61.633	49.360	10.951	1.00 15.47	SOLV
ATOM	42	OH2 WAT	342	89.589	43.811	5.959	1.00 18.08	SOLV
ATOM	43	OH2 WAT	343	70.742	35.952	14.932	1.00 34.03	SOLV
		OH2 WAT	344	89.836		26.657	1.00 18.11	
ATOM	44		_		28.590			SOLV
ATOM	45	OH2 WAT	345	70.822	32.764	1.461	1.00 22.35	SOLV
MOTA	46	OH2 WAT	346	63.056	34.653	0.491	1.00 29.51	SOLV
ATOM	47	OH2 WAT	347	58.054	46.282	2.363	1.00 10.00	SOLV
ATOM	48	OH2 WAT	348	67.914	58.660	-6.267	1.00 18.30	SOLV
ATOM	. 49	OH2 WAT	349	70.170	56.725	0.575	1.00 11.89	SOLV
ATOM	50	OH2 WAT	350	55.922	73.897			
						0.623	1.00 18.86	SOLV
MOTA	51	OH2 WAT	351	73.489	53.195	2.061	1.00 24.35	SOLV
ATOM	52	OH2 WAT	352	58.033	50.530	19.075	1.00 25.52	SOLV
MOTA	53	OH2 WAT	353	63.245	57.302	17.340	1.00 13.88	SOLV
MOTA	54	OH2 WAT	354	58.442	71.334	-5.670	1.00 17.51	SOLV
ATOM	55	OH2 WAT	355	62.535	61.154	16.706	1.00 12.38	SOLV
ATOM	56	OH2 WAT	356	66.949		-10.284	1.00 17.92	
ATOM	57	OH2 WAT	357	57.588	54.191	9.850		SOLV
							1.00 17.88	SOLV
MOTA	58	OH2 WAT	358	64.836	48.085	4.627	1.00 17.80	SOLV
MOTA	59	OH2 WAT	359	66.445	61.785	19.640	1.00 24.12	SOLV
ATOM	60	OH2 WAT	360	55.740	42.557	0.533	1.00 27.32	SOLV
ATOM	61	OH2 WAT	361	74.075	57.146	13.179	1.00 18.01	SOLV
ATOM	62	OH2 WAT	362	46.987	69.315	-2.545	1.00 11.87	SOLV
ATOM	63	OH2 WAT	363	53.842	52.266	-2.612	1.00 25.20	SOLV
MOTA	64	OH2 WAT	364	33.425	65.313	-4.686	1.00 28.97	
ATOM								sorv
	65	OH2 WAT	365	45.633	51.173	10.502	1.00 31.97	SOLV
MOTA	66	OH2 WAT	366	39.040	71.050	-0.722	1.00 20.81	SOLV
MOTA	67	OH2 WAT	367	54.517	67.335	-6.251	1.00 46.24	SOLV
MOTA	68	OH2 WAT	368	45.083	67.138	20.314	1.00 29.47	SOLV
ATOM	69	OH2 WAT	369	65.758	67.669	-6.655	1.00 14.69	SOLV
MOTA	70	OH2 WAT	370	44.943	78.174	12.948	1.00 23.88	SOLV
ATOM	71	OH2 WAT	371	37.141	57.403	1.723	1.00 23.72	
		OH2 WAT						SOLV
MOTA	72		372	62.407	66.806	13.368	1.00 13.36	SOLV
MOTA	73	OH2 WAT	373	50.776	47.263	5.661	1.00 38.22	SOLV
ATOM	74	OH2 WAT	374	56.697	47.264	11.752	1.00 24.75	SOLV
ATOM	75	OH2 WAT	375	42.566	60.884	15.739	1.00 16.25	SOLV
ATOM	76	OH2 WAT	376	59.299	74.342	13.838	1.00 31.27	SOLV
MOTA	77	OH2 WAT	377	72.976	63.691	-0.667	1.00 20.36	SOLV
ATOM	78	OH2 WAT	378	72.876	60.516	-6.752		
							1.00 34.24	SOLV
MOTA	79	OH2 WAT	379	63.998	68.760	16.371	1.00 19.04	SOLV
MOTA	80	OH2 WAT	380	44.947	66.728	-2.566	1.00 29.51	SOLV
MOTA	81	OH2 WAT	381	57.690	61.926	-9.414	1.00 29.01	SOLV
ATOM	82	OH2 WAT	382	44.595	80.810	5.831	1.00 27.43	SOLV
ATOM	83	OH2 WAT	383	78.065	36.583	24.121	1.00 14.08	SOLV
ATOM .		OH2 WAT	384	42.289	64.651	-0.868	1.00 25.57	SOLV
ATOM	85	OH2 WAT	385	59.851		~12.381	1.00 30.18	
ATOM		OH2 WAT						SOLV
	86		386	53.784	72.644	-4.782	1.00 22.35	SOLV
MOTA	· 87	OH2 WAT	387	72.793	27.922	8.925	1.00 32.13	SOLV
ATOM	88	OH2 WAT	388	57.224	68.062	-6.072	1.00 17.87	SOLV
MOTA	89	OH2 WAT	389	45.210	44.988	4.285	1.00 25.10	SOLV
ATOM	90	OH2 WAT	390	49.413	53.782	1.546	1.00 21.68	SOLV
ATOM	91	OH2 WAT	391	45.232	59.677	1.393	1.00 19.25	SOLV
MOTA	92	OH2 WAT	392	42.551	59.954	5.056	1.00 27.30	SOLV
ATOM	93	OH2 WAT	393	58.412	43.750	3.948	1.00 58.70	
ATOM	94	OH2 WAT	394	56.942	54.199		1.00 38.70	SOLV
						~2.588		SOLV
MOTA	95	OH2 WAT	395	55.216	51.994	9.824	1.00 13.25	SOLV
MOTA	96	OH2 WAT	396	51.642	54.651	14.874	1.00 10.00	SOLV
ATOM	97	OH2 WAT	397	48.690	56.156	13.991	1.00 28.59	SOLV
MOTA	98	OH2 WAT	398	74.412	37.913	0.396	1.00 12.55	SOLV
ATOM	99	OH2 WAT	399	81.920	53.968	18.267	1.00 14.05	SOLV
MOTA	100	OH2 WAT	400	70.413	41.780	1.170	1.00 16.68	SOLV
ATOM	101	OH2 WAT	401	71.098		2.407		
					53.544		1.00 27.63	SOLV
MOTA	102	OH2 WAT	402	94.383	32.979	9.497	1.00 27.97	SOLV
MOTA	103	OH2 WAT	403	70.765	66.069	16.389	1.00 38.09	SOLV
ATOM	104	OH2 WAT	404	78.651	34.890	29.495	1.00 48.60	SOLV
ATOM	105	OH2 WAT	405	80.289	39.811	24.727	1.00 20.74	SOLV
MOTA	106	OH2 WAT	406	63.627	47.414	7.301	1.00 40.21	SOLV
MOTA	107	OH2 WAT	407	74.679	30.772	11.524	1.00 37.03	
ATOM	108	OH2 WAT	408					SOLV
ATOM				80.240	36.041	26.681	1.00 27.42	SOLV
	109	OH2 WAT	409	84.971	25.909	18.426	1.00 24.96	SOLV
MOTA	110	OH2 WAT	410	57.832	41.294	5.792	1.00 71.90	SOLV
MOTA	111	OH2 WAT	411	55.484	68.139	-9.086	1.00 48.47	SOLV
MOTA	112	OH2 WAT	412	65.535	68.260	2.400	1.00 26.24	SOLV
MOTA	113	OH2 WAT	413	80.085	42.291	-3.144	1.00 26.49	SOLV
MOTA	114	OH2 WAT	414	82.088	37.456	27.733	1.00 42.54	SOLV
ATOM	115	OH2 WAT	415	61.020	53.195	21.566	1.00 38.16	SOLV
ATOM	116	OH2 WAT	416					
				55.968	70.365	-5.096	1.00 28.42	SOLV
MOTA	117	OH2 WAT	417	51.619	57.620	-0.487	1.00 41.81	SOLV

FIG. 6

# Compound C

FIG. 7



Compound E

**SUBSTITUTE SHEET (RULE 26)** 

a punodux

FIG. 8

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/05150

A. *CLASSIFICATION OF SUBJECT MATTER			
IPC(7) :G01N 9/00, 33/48 US CL :435/183: 702/22			
According to International Patent Classification (IPC) or to both national classification and IPC			
B. FIELDS SEARCHED			
Minimum documentation searched (classification system followed by classification symbols)			
U.S. : 435/183; 702/22			
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched			
NONE			
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)			
STN: WEST			
C. DOCUMENTS CONSIDERED TO BE RELEVANT			
Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.			
	-		
X	GOMIS-RUTH, F.X. et al. The helping hand of collagenase-3 8-14		
	(MMP-13: 2.7, ANG> crystal structure of its C-terminal haemopexin-like domain. Journal Mol. Biol. 1996, Vol. 264, No.		
	3, pages 556-566, see entire document		
X	US 6,008,243 A (BENDER et al.) 28 December 1999(28.12.99), see   1-7, 15-20		
	entire document.		
l			,
	<u> </u>		
Further documents are listed in the continuation of Box C. See patent family annex.			
Special categories of cited documents:  "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand		ication but cited to understand	
	ocument defining the general state of the art which is not considered be of particular relevance	*X" document of particular relevance; the	
	rlier document published on or after the international filing date	considered novel or cannot be consid	
cit	recument which may throw doubts on priority claim(s) or which is ted to establish the publication date of another citation or other secial reason (as specified)	"Y" document of particular relevance; th	e claimed invention cannot be
*O* do	ocument referring to an oral disclosure, use, exhibition or other	considered to involve an inventive combined with one or more other such being obvious to a person skilled in t	documents, such combination
*P* do	ocument published prior to the international filing date but later than	"&" document member of the same patent family	
Date of the actual completion of the international search  Date of mailing of the international search report			
12 JULY 2001		3,0 JUL 2007"	
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks  Authorized difficult  Authorize			
Commissioner of Patents and Trademarks Box PCT Washington D.C. 20231		AMY J. HARTTER	TUN
Washington, D.C. 20231  Facsimile No. (703) 305-3230		Telephone No. (703) 308-0196	

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/05150

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)			
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:			
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:			
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:			
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).			
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)			
This International Searching Authority found multiple inventions in this international application, as follows:			
Please See Extra Sheet.			
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.			
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:			
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:			
Remark on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.			

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/05150

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for as inventions to searched the appropriate search fees must be paid.

Group I which consists of claims 1-7 is distinct as it addresses itself to the solution complex of the mixture of MMP-13 and the defined "Compound A." The solution is clearly distinct and different from the crystal complex, active site and methods that are claimed in succeeding groups and according claims.

Group II consists of claims 8-14. These claims pertain to the actual product of the crystal complexion its entirety. Thus it is distinct from Groups I and Groups 3-4. The group claims the whole crystal known as "Compound A". and the crystal is not in any other type of alternate environment or with any additional accountements.

Group III encompasses the claims of 15-20. These claims consist of the active site of the molecule of MMP-13. This chemical is a portion of the solution claimed in the first group and thus separate and distinct from the solution of Group I or the separate entity of "Compound A" that is claimed in Group 2. Thus these Groups are separate.

Group IV consists of claims 21-32 which claim a method of identifying an inhibitor or activator of the MMP-13 compound. The method that is embodied within this Group is clearly different from the proceeding groups. Firstly the claims within Group 4 are directed toward a method of accomplishing the task of identifying different entities and not a product itself. Secondly its actions are addressed to entities outside the compound itself and not limited to "Compound A" of the MMP-13. Based on the aforementioned reasons and the distinct nature of the claims defined in each of the groups, the instant application has a lack of unity due to each group having a different Special Technical Feature a summarized above for each group.